Package ‘rstan’

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Description User-facing R functions are provided to parse, compile, test, estimate, and analyze Stan models by accessing the header-only Stan library provided by the ‘StanHeaders’ package. The Stan project develops a probabilistic programming language that implements full Bayesian statistical inference via Markov Chain Monte Carlo, rough Bayesian inference via ‘variational’ approximation, and (optionally penalized) maximum likelihood estimation via optimization. In all three cases, automatic differentiation is used to quickly and accurately evaluate gradients without burdening the user with the need to derive the partial derivatives.

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BugReports https://github.com/stan-dev/rstan/issues/

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Description

RStan is the R interface to the Stan C++ package. The RStan interface (rstan R package) provides:

- Full Bayesian inference using the No-U-Turn sampler (NUTS), a variant of Hamiltonian Monte Carlo (HMC)
- Approximate Bayesian inference using automatic differentiation variational inference (ADVI)
- Penalized maximum likelihood estimation using L-BFGS optimization

For documentation on Stan itself, including the manual and user guide for the modeling language, case studies and worked examples, and other tutorial information visit the Users section of the Stan website:

- mc-stan.org/users/documentation

Other R packages from the Stan Development Team

Various related R packages are also available from the Stan Development Team:

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There are also many other important contributors to RStan (github.com/rstan). Please use ‘Stan Development Team’ whenever citing the R interface to Stan. A BibTex entry is available from http://mc-stan.org/rstan/authors or citation("rstan").

See Also

- The RStan vignettes: http://mc-stan.org/rstan/articles/.
- `stan` for details on fitting models and `stanfit` for information on the fitted model objects.
- The `lookup` for finding a function in the Stan language that corresponds to a R function or name.
- https://github.com/stan-dev/rstan/issues/ to submit a bug report or feature request.

Examples

```r
# Not run:

stammodelcode <- "
data {
  int<lower=0> N;
  real y[N];
}

parameters {
  real mu;
}

model {
  target += normal_lpdf(mu | 0, 10);
  target += normal_lpdf(y | mu, 1);
}
"

y <- rnorm(20)
dat <- list(N = 20, y = y);
fit <- stan(model_code = stammodelcode, model_name = "example",
            data = dat, iter = 2012, chains = 3, verbose = TRUE,
            sample_file = file.path(tempdir(), 'norm.csv'))
print(fit)

# extract samples
e <- extract(fit, permuted = FALSE) # return a list of arrays
str(e)

arr <- as.array(fit) # return an array
str(arr)

mat <- as.matrix(fit) # return a matrix
```

as.array

str(mat)

## End(Not run)

---

as.array Create array, matrix, or data.frame objects from samples in a stanfit object

**Description**

The samples (without warmup) included in a stanfit object can be coerced to an array, matrix, or data.frame. Methods are also provided for checking and setting names and dimnames.

**Usage**

```r
## S3 method for class 'stanfit'
as.array(x, ...)
## S3 method for class 'stanfit'
as.matrix(x, ...)
## S3 method for class 'stanfit'
as.data.frame(x, ...)
## S3 method for class 'stanfit'
is.array(x)
## S3 method for class 'stanfit'
dim(x)
## S3 method for class 'stanfit'
dimnames(x)
## S3 method for class 'stanfit'
names(x)
## S3 replacement method for class 'stanfit'
names(x) <- value
```

**Arguments**

- `x` An object of S4 class stanfit.
- `...` Additional parameters that can be passed to `extract` for extracting samples from x. For now, pars is the only additional parameter supported.
- `value` For the names replacement method, a character vector to set/replace the parameter names in x.

**Details**

as.array and as.matrix can be applied to a stanfit object to coerce the samples without warmup to array or matrix. The as.data.frame method first calls as.matrix and then coerces this matrix to a data.frame.

The array has three named dimensions: iterations, chains, parameters. For as.matrix, all chains are combined, leaving a matrix of iterations by parameters.
Value

as.array, as.matrix, and as.data.frame return an array, matrix, and data.frame, respectively. dim and dimnames return the dim and dimnames of the array object that could be created, while names returns the third element of the dimnames, which are the names of the margins of the posterior distribution. The names assignment method allows for assigning more interpretable names to them. is.array returns TRUE for stanfit objects that include samples; otherwise FALSE.

When the stanfit object does not contain samples, empty objects are returned from as.array, as.matrix, as.data.frame, dim, dimnames, and names.

See Also

S4 class stanfit and its method extract

Examples

```r
## Not run:
ex_model_code <- ' parameters {
  real alpha[2,3];
  real beta[2];
}
model {
  for (i in 1:2) for (j in 1:3)
    alpha[i, j] ~ normal(0, 1);
  for (i in 1:2)
    beta[i] ~ normal(0, 2);
  # beta ~ normal(0, 2) // vectorized version
}

## fit the model
fit <- stan(model_code = ex_model_code, chains = 4)
dim(fit)
dimnames(fit)
is.array(fit)
a <- as.array(fit)
m <- as.matrix(fit)
d <- as.data.frame(fit)

## End(Not run)
```

As.mcmc.list

Create an mcmc.list from a stanfit object

Description

Create an mcmc.list (coda) from a stanfit object.
check_hmc_diagnostics

Usage
As.mcmc.list(object, pars, include = TRUE, ...)

Arguments

object  object of class "stanfit"
pars    optional character vector of parameters to include
include logical scalar indicating whether to include (the default) or exclude the parameters named in pars
...

Value
An object of class mcmc.list.

Description
These functions print summaries of important HMC diagnostics or extract those diagnostics from a stanfit object. See the Details section, below.

Usage
check_hmc_diagnostics(object)
check_divergences(object)
check_treedepth(object)
check_energy(object)
get_divergent_iterations(object)
get_max_treedepth_iterations(object)
get_num_leapfrog_per_iteration(object)
get_num_divergent(object)
get_num_max_treedepth(object)
get_bfmi(object)
get_low_bfmi_chains(object)

Arguments

object A stanfit object.
Details

The check_hmc_diagnostics function calls the other check_* functions internally and prints an overall summary, but the other functions can also be called directly:

- check_divergences prints the number (and percentage) of iterations that ended with a divergence,
- check_treedepth prints the number (and percentage) of iterations that saturated the max treedepth,
- check_energy prints E-BFMI values for each chain for which E-BFMI is less than 0.2.

The get_* functions are for programmatic access to the diagnostics.

- get_divergent_iterations and get_max_treedepth_iterations return a logical vector indicating problems for individual iterations,
- get_num_divergences and get_num_max_treedepth return the number of offending iterations,
- get_num_leapfrog_per_iteration returns an integer vector with the number of leapfrog evaluations for each iteration,
- get_bfmi returns per-chain E-BFMI values and get_low_bfmi_chains returns the indices of chains with low E-BFMI.

The following are several of many resources that provide more information on these diagnostics:

- Brief explanations of some of the problems detected by these diagnostics can be found in the Brief Guide to Stan’s Warnings.
- Betancourt (2017) provides much more depth on these diagnostics as well as a conceptual introduction to Hamiltonian Monte Carlo in general.
- Gabry et al. (2018) and the bayesplot package vignettes demonstrate various visualizations of these diagnostics that can be made in R.

References


Examples

```r
## Not run:
schools <- stan_demo("eight_schools")
check_hmc_diagnostics(schools)
check_divergences(schools)
check_treedepth(schools)
```
Diagnostic plots

```r
check_energy(schools)
```

## End(Not run)

### RStan Diagnostic plots

#### Description

Diagnostic plots for HMC and NUTS using ggplot2.

#### Usage

```r
stan_diag(object, information = c("sample", "stepsize", "treedepth", "divergence"),
          chain = 0, ...)
stan_par(object, par, chain = 0, ...)
stan_rhat(object, par, chain = 0, ...)
stan_ess(object, par, chain = 0, ...)
stan_mcse(object, par, chain = 0, ...)
```

#### Arguments

- **object**: A stanfit or stanreg object.
- **information**: The information to be contained in the diagnostic plot.
- **par, pars**: The name of a single scalar parameter (par) or one or more parameter names (pars).
- **chain**: If chain = 0 (the default) all chains are combined. Otherwise the plot for chain is overlaid on the plot for all chains combined.
- **...**: For `stan_diag` and `stan_par`, optional arguments to `arrangeGrob`. For `stan_rhat`, `stan_ess`, and `stan_mcse`, optional arguments to `stat_bin`.

#### Details

- **`stan_rhat`, `stan_ess`, `stan_mcse`**: Respectively, these plots show the distribution of the Rhat statistic, the ratio of effective sample size to total sample size, and the ratio of Monte Carlo standard error to posterior standard deviation for the estimated parameters. These plots are not intended to identify individual parameters, but rather to allow for quickly identifying if the estimated values of these quantities are desirable for all parameters.
- **`stan_par`**: Calling `stan_par` generates three plots: (i) a scatterplot of par vs. the accumulated log-posterior (lp__), (ii) a scatterplot of par vs. the average Metropolis acceptance rate (accept_stat), and (iii) a violin plot showing the distribution of par at each of the sampled step sizes (one per chain). For the scatterplots, red points are superimposed to indicate which (if any) iterations encountered a divergent transition. Yellow points indicate a transition that hit the maximum treedepth rather than terminated its evolution normally.
expose_stan_functions

The information argument is used to specify which plots stan_diag should generate:

- `information='sample'` Histograms of lp__ and accept_stat, as well as a scatterplot showing their joint distribution.
- `information='stepsizes'` Violin plots showing the distributions of lp__ and accept_stat at each of the sampled step sizes (one per chain).
- `information='treedepth'` Histogram of treedepth and violin plots showing the distributions of lp__ and accept_stat for each value of treedepth.
- `information='divergence'` Violin plots showing the distributions of lp__ and accept_stat for iterations that encountered divergent transitions (divergent=1) and those that did not (divergent=0).

Value

For `stan_diag` and `stan_par`, a list containing the ggplot objects for each of the displayed plots. For `stan_rhat`, `stan_ess`, and `stan_mcse`, a single ggplot object.

Note


See Also

List of RStan plotting functions, Plot options

Examples

```r
## Not run:
fit <- stan_demo("eight_schools")
stan_diag(fit, info = 'sample')  # shows three plots together
samp_info <- stan_diag(fit, info = 'sample')  # saves the three plots in a list
samp_info[3]  # access just the third plot
stan_diag(fit, info = 'sample', chain = 1)  # overlay chain 1
stan_par(fit, par = "mu")
## (Not run)
```

Description

The Stan modeling language allows users to define their own functions in a functions block at the top of a Stan program. The `expose_stan_functions` utility function uses `sourceCpp` to export those user-defined functions to the specified environment for testing inside R or for doing posterior predictive simulations in R rather than in the generated quantities block of a Stan program.
**Usage**

```r
expose_stan_functions(stanmodel, includes = NULL, 
                        show_compiler_warnings = FALSE, ...)
get_rng(seed = 0L)
get_stream()
```

**Arguments**

- **stanmodel**: A `stanmodel` object, a `stanfit` object, a list produced by `stanc` or the path to a Stan program (.stan file). In any of these cases, the underlying Stan program should contain a non-empty functions block.
- **includes**: If not `NULL` (the default), then a character vector of length one (possibly containing one or more `"\n"`) of the form `"#include "/full/path/to/my_header.hpp"`, which will be inserted into the C++ code in the model's namespace and can be used to provide definitions of functions that are declared but not defined in `stanmodel`.
- **show_compiler_warnings**: Logical scalar defaulting to `FALSE` that controls whether compiler warnings, which can be numerous and have never been relevant, are shown.
- **seed**: An integer vector of length one indicating the state of Stan's pseudo-random number generator.
- **...**: Further arguments passed to `sourceCpp`.

**Details**

The `expose_stan_functions` function requires as much compliance with the C++14 standard as is implemented in the RTools toolchain for Windows. On Windows, you will likely need to specify `CXX14 = g++ -std=c++1y` in the file whose path is `normalizePath("~/.R/Makevars")` in order for `expose_stan_functions` to work. Outside of Windows, the necessary compiler flags are set programatically, which is likely to suffice.

There are a few special types of user-defined Stan functions for which some additional details are relevant:

**(P)RNG functions**: If a user-defined Stan function ends in `_rng`, then it can use the Boost pseudo-random number generator used by Stan. When exposing such functions to R, `base_rng__` and `pstream__` arguments will be added to the formals. The `base_rng__` argument should be passed the result of a call to `get_rng` (perhaps specifying its `seed` argument for reproducibility) and the `pstream__` should be passed the result of a call to `get_stream`, which can be used to see the result of `print` and `reject` calls in the user-defined Stan functions. These arguments default to `get_stream()` and `get_rng()` respectively.

**LP functions**: If a user-defined Stan function ends in `_lp`, then it can modify the log-probability used by Stan to evaluate Metropolis proposals or as an objective function for optimization. When exposing such functions to R, a `lp__` argument will be added to the formals. This `lp__` argument defaults to zero, but a `double` precision scalar may be passed to this argument when the function is called from R. Such a user-defined Stan function can terminate with `return target();` or can execute `print(target());` to verify that the calculation is correct.
Value

The names of the new functions in env are returned invisibly.

See Also

sourceCpp and the section in the Stan User Manual on user-defined functions

Examples

```r
## Not run:
model_code <-
',
functions {
  real standard_normal_rng() {
    return normal_rng(0,1);
  }
},

expose_stan_functions(stanc(model_code = model_code))
standard_normal_rng()
PRNG <- get_rng(seed = 3)
o <- get_stream()
standard_normal_rng(PRNG, o)

## End(Not run)
```

---

extract | Extract samples from a fitted Stan model

Description

Extract samples from a fitted model represented by an instance of class stanfit.

Usage

```r
## S4 method for signature 'stanfit'
xextract(object, pars, permuted = TRUE, inc_warmup = FALSE, 
         include = TRUE)
```

Arguments

- **object**: An object of class stanfit.
- **pars**: An optional character vector providing the parameter names (or other quantity names) of interest. If not specified, all parameters and other quantities are used. The log-posterior with name lp__ is also included by default.
- **permuted**: A logical scalar indicating whether the draws after the warmup period in each chain should be permuted and merged. If FALSE, the original order is kept. For each stanfit object, the permutation is fixed (i.e., extracting samples a second time will give the same sequence of iterations).
inc_warmup A logical scalar indicating whether to include the warmup draws. This argument is only relevant if permuted is FALSE.

include A logical scalar indicating whether the parameters named in pars should be included (TRUE) or excluded (FALSE).

Value

When permuted = TRUE, this function returns a named list, every element of which is an array representing samples for a parameter with all chains merged together.

When permuted = FALSE, an array is returned; the first dimension is for the iterations, the second for the number of chains, the third for the parameters. Vectors and arrays are expanded to one parameter (a scalar) per cell, with names indicating the third dimension. See the examples (with comments) below. The monitor function can be applied to the returned array to obtain a summary (similar to the print method for stanfit objects).

Methods

extract signature(object = "stanfit") Extract samples from a fitted model represented by an instance of class stanfit.

See Also

S4 class stanfit, as.array.stanfit, and monitor

Examples

```r
## Not run:
ex_model_code <- 'parameters {
  real alpha[2,3];
  real beta[2];
}
model {
  for (i in 1:2) for (j in 1:3)
    alpha[i, j] ~ normal(0, 1);
  for (i in 1:2)
    beta ~ normal(0, 2);
}
'

## fit the model
fit <- stan(model_code = ex_model_code, chains = 4)

## extract alpha and beta with 'permuted = TRUE'
fit_ss <- extract(fit, permuted = TRUE) # fit_ss is a list
## list fit_ss should have elements with name 'alpha', 'beta', 'lp__'
alpha <- fit_ss$alpha
beta <- fit_ss$beta
## or extract alpha by just specifying pars = 'alpha'
alpha2 <- extract(fit, pars = 'alpha', permuted = TRUE)$alpha
print(identical(alpha, alpha2))
```
## or extract alpha by excluding beta and lp_

alpha3 <- extract(fit, pars = c('beta', 'lp_'),
                 permuted = TRUE, include = FALSE)$alpha
print(identical(alpha, alpha3))

## get the samples for alpha[1,1] and beta[2]
alpha_11 <- alpha[, 1, 1]
beta_2 <- beta[, 2]

## extract samples with 'permuted = FALSE'
fit_ss2 <- extract(fit, permuted = FALSE) # fit_ss2 is an array

## the dimensions of fit_ss2 should be
## "# of iterations * # of chains * # of parameters"
dim(fit_ss2)

## since the third dimension of 'fit_ss2' indicates
## parameters, the names should be
## alpha[1,1], alpha[2,1], alpha[1,2], alpha[2,2],
## alpha[1,3], alpha[2,3], beta[1], beta[2], and lp_
## 'lp_' (the log-posterior) is always included
## in the samples.
dimnames(fit_ss2)

## End(Not run)

# Create a stanfit object from reading CSV files of samples (saved in rstan
# package) generated by function stan for demonstration purpose from model as follows.
#
excode <- 'transformed data {
  real y[20];
  y[1] <- 0.5796; y[2] <- 0.2276; y[3] <- -0.2959;
  y[13] <- 0.5215; y[14] <- -1.6044; y[15] <- -0.6783;
  y[16] <- 0.9459; y[17] <- -0.382; y[18] <- 0.7619;
  y[19] <- 0.1006; y[20] <- -1.7461;
}
parameters {
  real mu;
  real<lower=0, upper=10> sigma;
  vector[2] z[3];
  real<lower=0> alpha;
}
model {
  y ~ normal(mu, sigma);
  for (i in 1:3)
    z[i] ~ normal(0, 1);
  alpha ~ exponential(2);
}
### extract_sparse_parts

`extract_sparse_parts` is used to extract the compressed representation of a sparse matrix.

#### Description

Create a list of vectors that represents a sparse matrix.

#### Usage

```r
extract_sparse_parts(A)
```

#### Arguments

- `A` (A matrix or Matrix)

#### Details

The Stan Math Library has a function called `csr_matrix_times_vector`, which inputs a matrix in compressed row storage form and a dense vector and returns their product without fillin. To use the `csr_matrix_times_vector` function with a large sparse matrix, it is optimal in terms of memory to simply pass the three vectors that characterize the compressed row storage form of the matrix to the data block of the Stan program. The `extract_sparse_parts` function provides a convenient means of obtaining these vectors.

#### Value

A named list with components

1. `w` A numeric vector containing the non-zero elements of `A`.
2. `v` An integer vector containing the column indices of the non-zero elements of `A`.
3. `u` An integer vector indicating where in `w` a given row’s non-zero values start.
Examples

```r
A <- rbind(
  c(1L, 2L, 0L, 0L),
  c(0L, 0L, 0L, 0L),
  c(0L, 0L, 0L, 5L),
  c(8L, 0L, 9L, 3L)
)
str(extract_sparse_parts(A))
```

---

**gqs**

*Draw samples of generated quantities from a Stan model*

Description

Draw samples from the generated quantities block of a `stanmodel`.

Usage

```r
## S4 method for signature 'stanmodel'
gqs(object, data = list(), draws, seed = sample.int(.Machine$integer.max, size = 1L))
```

Arguments

- **object**: An object of class `stanmodel`.
- **data**: A named list or environment providing the data for the model or a character vector for all the names of objects used as data. See the **Passing data to Stan** section in `stan`.
- **draws**: A matrix of posterior draws, typically created by calling `as.matrix` on a `stanfit`.
- **seed**: The seed for random number generation. The default is generated from 1 to the maximum integer supported by `R` on the machine. When a seed is specified by a number, `as.integer` will be applied to it. If `as.integer` produces `NA`, the seed is generated randomly. The seed can also be specified as a character string of digits, such as "12345", which is converted to integer.

Value

An object of S4 class `stanmodel` representing the fitted results.

Methods

- `object signature(object = "stanmodel")` Evaluate the generated quantities block of a Stan program by supplying data and the draws output from a previous Stan program.

See Also

`stanmodel, stanfit, stan`
Examples

```r
## Not run:
m <- stan_model(model_code = 'parameters {real y;}
model {y ~ normal(0,1);}')
f <- sampling(m, iter = 300)
mc <-
  parameters {real y;}
generated quantities {real y_rep = normal_rng(y, 1);}
'm2 <- stan_model(model_code = mc)
f2 <- gqs(m2, draws = as.matrix(f))
f2

## End(Not run)
```

Description

Using model’s `log_prob` and `grad_log_prob` take values from the unconstrained space of model parameters and (by default) return values in the same space. Sometimes we need to convert the values of parameters from their support defined in the parameters block (which might be constrained, and for simplicity, we call it the constrained space) to the unconstrained space and vice versa. The `constrain_pars` and `unconstrain_pars` functions are used for this purpose.

Usage

```r
## S4 method for signature 'stanfit'
log_prob(object, upars, adjust_transform = TRUE, gradient = FALSE)

## S4 method for signature 'stanfit'
grad_log_prob(object, upars, adjust_transform = TRUE)

## S4 method for signature 'stanfit'
get_num_upars(object)

## S4 method for signature 'stanfit'
constrain_pars(object, upars)

## S4 method for signature 'stanfit'
unconstrain_pars(object, pars)
```

Arguments

- `object` An object of class `stanfit`.
pars  An list specifying the values for all parameters on the constrained space.
upars  A numeric vector for specifying the values for all parameters on the unconstrained space.
adjust_transform  Logical to indicate whether to adjust the log density since Stan transforms parameters to unconstrained space if it is in constrained space. Set to FALSE to make the function return the same values as Stan’s lp__ output.
gradient  Logical to indicate whether gradients are also computed as well as the log density.

Details
Stan requires that parameters be defined along with their support. For example, for a variance parameter, we must define it on the positive real line. But inside Stan’s samplers all parameters defined on the constrained space are transformed to an unconstrained space amenable to Hamiltonian Monte Carlo. Because of this, Stan adjusts the log density function by adding the log absolute value of the Jacobian determinant. Once a new iteration is drawn, Stan transforms the parameters back to the original constrained space without requiring interference from the user. However, when using the log density function for a model exposed to R, we need to be careful. For example, if we are interested in finding the mode of parameters on the constrained space, we then do not need the adjustment. For this reason, the log_prob and grad_log_prob functions accept an adjust_transform argument.

Value
log_prob returns a value (up to an additive constant) the log posterior. If gradient is TRUE, the gradients are also returned as an attribute with name gradient.
grad_log_prob returns a vector of the gradients. Additionally, the vector has an attribute named log_prob being the value the same as log_prob is called for the input parameters.
get_num_upars returns the number of parameters on the unconstrained space.
constrain_pars returns a list and unconstrain_pars returns a vector.

Methods
log_prob signature(object = "stanfit") Compute lp__, the log posterior (up to an additive constant) for the model represented by a stanfit object. Note that, by default, log_prob returns the log posterior in the unconstrained space Stan works in internally. set adjust_transform = FALSE to make the values match Stan’s output.
grad_log_prob signature(object = "stanfit") Compute the gradients for log_prob as well as the log posterior. The latter is returned as an attribute.
get_num_upars signature(object = "stanfit") Get the number of unconstrained parameters.
constrain_pars signature(object = "stanfit") Convert values of the parameter from unconstrained space (given as a vector) to their constrained space (returned as a named list).
unconstrain_pars signature(object = "stanfit") Contrary to constrained, convert values of the parameters from constrained to unconstrained space.
### loo.stanfit

**Approximate leave-one-out cross-validation**

**Description**

A `loo` method that is customized for `stanfit` objects. The `loo` method for `stanfit` objects—a wrapper around the `loo.array` (`loo` package)—computes PSIS-LOO CV, approximate leave-one-out cross-validation using Pareto smoothed importance sampling (Vehtari, Gelman, and Gabry, 2017a, 2017b).
Usage

```r
## S3 method for class 'stanfit'
loo(x, pars = "log_lik",
   ..., save_psis = FALSE,
   cores = getOption("mc.cores", 1))
```

Arguments

- `x`: An object of S4 class `stanfit`.
- `pars`: Name of transformed parameter or generated quantity in the Stan program corresponding to the pointwise log-likelihood. If not specified the default behavior is to look for "log_lik".
- `cores`: Number of cores to use for parallelization. The default is 1 unless `cores` is specified or the `mc.cores` option has been set.
- `save_psis`: Should the intermediate results from `psis` be saved in the returned object? The default is `FALSE`. This can be useful to avoid repeated computation when using other functions in the `loo` and `bayesplot` packages.
- `...`: Ignored.

Details

Stan does not automatically compute and store the log-likelihood. It is up to the user to incorporate it into the Stan program if it is to be extracted after fitting the model. In a Stan program, the pointwise log likelihood can be coded as a vector in the transformed parameters block (and then summed up in the model block) or it can be coded entirely in the generated quantities block. We recommend using the generated quantities block so that the computations are carried out only once per iteration rather than once per HMC leapfrog step.

For example, the following is the generated quantities block for computing and saving the log-likelihood for a linear regression model with `n` data points, outcome `y`, predictor matrix `X` (including column of 1s for intercept), coefficients `beta`, and standard deviation `sigma`:

```stan
vector[N] log_lik;
for (n in 1:N) log_lik[n] = normal_lpdf(y[n] | X[n, ] * beta, sigma);
```

Value

A list with class `c(psis_loo, "loo")`, as detailed in the `loo.array` documentation.

References


See Also

- The loo package documentation, including the vignettes for many examples (http://mc-stan.org/loo).
- loo_model_weights for model averaging/weighting via stacking or pseudo-BMA weighting.

Examples

```r
## Not run:
# Generate a dataset from N(0,1)
N <- 100
y <- rnorm(N, 0, 1)

# Suppose we have three models for y:
# 1) y ~ N(-1, sigma)
# 2) y ~ N(0.5, sigma)
# 3) y ~ N(0.6, sigma)
#
# stan_code <- "
data {
    int N;
    vector[N] y;
    real mu_fixed;
}

parameters {
    real<lower=0> sigma;
}

model {
    sigma ~ exponential(1);
    y ~ normal(mu_fixed, sigma);
}

generated quantities {
    vector[N] log_lik;
    for (n in 1:N) log_lik[n] = normal_lpdf(y[n] | mu_fixed, sigma);
}"

mod <- stan_model(model_code = stan_code)
fit1 <- sampling(mod, data=list(N=N, y=y, mu_fixed=-1))
fit2 <- sampling(mod, data=list(N=N, y=y, mu_fixed=0.5))
fit3 <- sampling(mod, data=list(N=N, y=y, mu_fixed=0.6))

# use the loo method for stanfit objects
loo1 <- loo(fit1, pars = "log_lik")
print(loo1)

# which is equivalent to
LL <- as.array(fit1, pars = "log_lik")
r_eff <- loo::relative_eff(exp(LL))
loo1b <- loo::loo.array(LL, r_eff = r_eff)
print(loo1b)

# compute loo for the other models
loo2 <- loo(fit2)
```

lookup <- loo(fit3)

# stacking weights
wts <- loo::loo_model_weights(list(loo1, loo2, loo3), method = "stacking")
print(wts)

## End(Not run)

```r
lookup
```

Look up the Stan function that corresponds to a R function or name.

Description

This function helps to map between R functions and Stan functions.

Usage

```r
lookup(FUN, ReturnType = character())
```

Arguments

- `FUN` A character string naming a R function or a R function for which the (near) equivalent Stan function is sought. If no matching R function is found, FUN is reinterpreted as a `regexp` and matches are sought.
- `ReturnType` A character string of positive length naming a valid return type for a Stan function: `int`, `int[]`, `matrix`, `real`, `real[,]`, `real[]`, `row_vector`, `T[]`, `vector`, or `void`. If "ANY" is passed, then the entire `data.frame` is returned and can be inspected with the `View` function, for example.

Value

Ordinarily, a `data.frame` with rows equal to the number of partial matches and four columns:

1. `StanFunction` Character string for the Stan function’s name.
2. `Arguments` Character string indicating the arguments to that Stan function.
3. `ReturnType` Character string indicating the return type of that Stan function.
4. `Page` Integer indicating the page of the Stan reference manual where that Stan function is defined.

If there are no matching Stan functions, a character string indicating so is returned.

References


Examples

lookup(dnorm)    # Stan equivalents for the normal PDF (in log form)
lookup("foo")   # fails
lookup("Student") # succeeds even though there is no such R function
lookup("*poisson") # every Stan function that starts with poisson

makeconf_path

Obtain the full path of file Makeconf

Description

Obtain the full path of file Makeconf, in which, for example the flags for compiling C/C++ code are configured.

Usage

makeconf_path()

Details

The configuration for compiling shared objects using R CMD SHLIB are set in file Makeconf. To change how the C++ code is compiled, modify this file. For RStan, package inline compiles the C++ code using R CMD SHLIB. To speed up compiled Stan models, increase the optimization level to -O3 defined in property CXXFLAGS in the file Makeconf. This file may also be modified to specify alternative C++ compilers, such as clang++ or later versions of g++.

Value

An character string for the full path of file Makeconf.

See Also

stan

Examples

makeconf_path()
monitor  Compute summaries of MCMC draws and monitor convergence

Description

Similar to the print method for stanfit objects, but monitor takes an array of simulations as its argument rather than a stanfit object. For a 3-D array (iterations * chains * parameters) of MCMC draws, monitor computes means, standard deviations, quantiles, Monte Carlo standard errors, split Rhats, and effective sample sizes. By default, half of the iterations are considered warmup and are excluded.

Usage

monitor(sims, warmup = floor(dim(sims)[1]/2),
        probs = c(0.025, 0.25, 0.5, 0.75, 0.975),
        digits_summary = 1, print = TRUE, ...)
## S3 method for class 'simsummary'
print(x, digits = 3, se = FALSE, ...)
## S3 method for class 'simsummary'
x[i, j, drop = if (missing(i)) TRUE else length(j) == 1]

Arguments

sims     A 3-D array (iterations * chains * parameters) of MCMC simulations from any MCMC algorithm.
warmup   The number of warmup iterations to be excluded when computing the summaries. The default is half of the total number of iterations. If sims doesn’t include the warmup iterations then warmup should be set to zero.
probs    A numeric vector specifying quantiles of interest. The defaults is c(0.025,0.25,0.5,0.75,0.975).
digits_summary The number of significant digits to use when printing the summary, defaulting to 1. Applies to the quantities other than the effective sample size, which is always rounded to the nearest integer.
print    Logical, indicating whether to print the summary after the computations are performed.
...      Additional arguments passed to the underlying print method.
x        An object of class simsummary created by monitor
digits  An integer scalar defaulting to 3 for the number of decimal places to print
se       A logical scalar defaulting to FALSE indicating whether to print the estimated standard errors of the estimates
i        A vector indicating which rows of the object created by monitor to select
j        A vector indicating which columns of the object created by monitor to select
drop    A logical scalar indicating whether the resulting object should return a vector where possible
optimizing

Value

A 2-D array with rows corresponding to parameters and columns to the summary statistics that can be printed and subset.

References


See Also

S4 class stanfit and particularly its print method.

Examples

csvfiles <- dir(system.file('misc', package = 'rstan'),
    pattern = 'rstan_doc_ex_[0-9].csv', full.names = TRUE)
fit <- read_stan_csv(csvfiles)
# The following is just for the purpose of giving an example
# since print can be used for a stanfit object.
monitor(extract(fit, permuted = FALSE, inc_warmup = TRUE))

optimizing

Obtain a point estimate by maximizing the joint posterior

Description

Obtain a point estimate by maximizing the joint posterior from the model defined by class stanmodel.

Usage

## S4 method for signature 'stanmodel'
optimizing(object, data = list(),
    seed = sample.int(.Machine$integer.max, 1), init = 'random',
    check_data = TRUE, sample_file = NULL,
    algorithm = c("LBFGS", "BFGS", "Newton"),
    verbose = FALSE, hessian = FALSE, as_vector = TRUE,
    draws = 0, constrained = TRUE, importance_resampling = FALSE, ...)

Arguments

object An object of class stanmodel.
data A named list or environment providing the data for the model or a character vector for all the names of objects used as data. See the Passing data to Stan section in stan.
seed

The seed for random number generation. The default is generated from 1 to the maximum integer supported by R on the machine. Even if multiple chains are used, only one seed is needed, with other chains having seeds derived from that of the first chain to avoid dependent samples. When a seed is specified by a number, as.integer will be applied to it. If as.integer produces NA, the seed is generated randomly. The seed can also be specified as a character string of digits, such as "12345", which is converted to integer.

init

Initial values specification. See the detailed documentation for the init argument in stan with one exception. If specifying inits using a list then only a single named list of values should be provided. For example, to initialize a parameter alpha to value1 and beta to value2 you can specify list(alpha = value1, beta = value2).

check_data

Logical, defaulting to TRUE. If TRUE the data will be preprocessed; otherwise not. See the Passing data to Stan section in stan.

sample_file

A character string of file name for specifying where to write samples for all parameters and other saved quantities. If not provided, files are not created. When the folder specified is not writable, tempdir() is used.

algorithm

One of "Newton", "BFGS", and "LBFGS" (the default) indicating which optimization algorithm to use.

verbose

TRUE or FALSE (the default): flag indicating whether to print intermediate output from Stan on the console, which might be helpful for model debugging.

hessian

TRUE or FALSE (the default): flag indicating whether to calculate the Hessian (via numeric differentiation of the gradient function in the unconstrained parameter space).

as_vector

TRUE (the default) or FALSE: flag indicating whether a vector is used to store the point estimate found. A list can be used instead by specifying it to be FALSE.

draws

A non-negative integer (that defaults to zero) indicating how many times to draw from a multivariate normal distribution whose parameters are the mean vector and the inverse negative Hessian in the unconstrained space.

constrained

A logical scalar indicating, if draws > 0, whether the draws should be transformed to the constrained space defined in the parameters block of the Stan program. Defaults to TRUE.

importance_resampling

A logical scalar (defaulting to FALSE) indicating whether to do importance resampling to compute diagnostics on the draws from the normal approximation to the posterior distribution.

... Other optional parameters:

- iter (integer), the maximum number of iterations, defaulting to 2000.
- save_iterations (logical), a flag indicating whether to save the iterations, defaulting to FALSE.
- refresh (integer), the number of iterations between screen updates, defaulting to 100.
- init_alpha (double), for BFGS and LBFGS, the line search step size for first iteration, defaulting to 0.001.
- tol_obj (double), for BFGS and LBFGS, the convergence tolerance on changes in objective function value, defaulting to 1e-12.
• `tol_rel_obj(double)`, for BFGS and LBFGS, the convergence tolerance on relative changes in objective function value, defaulting to 1e4.
• `tol_grad(double)`, for BFGS and LBFGS, the convergence tolerance on the norm of the gradient, defaulting to 1e-8.
• `tol_rel_grad(double)`, for BFGS and LBFGS, the convergence tolerance on the relative norm of the gradient, defaulting to 1e7.
• `tol_param(double)`, for BFGS and LBFGS, the convergence tolerance on changes in parameter value, defaulting to 1e-8.
• `history_size(integer)`, for LBFGS, the number of update vectors to use in Hessian approximations, defaulting to 5.

Refer to the manuals for both CmdStan and Stan for more details.

Value

```r
par
```
The point estimate found. Its form (vector or list) is determined by the `as_vector` argument.

```r
value
```
The value of the log-posterior (up to an additive constant, the "lp__" in Stan) corresponding to `par`.

```r
return_code
```
The value of the return code from the optimizer; anything that is not zero is problematic.

```r
hessian
```
The Hessian matrix if `hessian` is `TRUE`.

```r
theta_tilde
```
If `draws > 0`, the matrix of parameter draws in the constrained or unconstrained space, depending on the value of the `constrained` argument.

```r
log_p
```
If `draws > 0`, a vector of length `draws` that contains the value of the log-posterior evaluated at each row of `theta_tilde`.

```r
log_g
```
If `draws > 0`, a vector of length `draws` that contains the value of the logarithm of the multivariate normal density evaluated at each row of `theta_tilde`.

If the optimization is not completed for reasons such as feeding wrong data, it returns NULL.

Methods

```r
optimizing signature(object = "stanmodel")
```
Call Stan’s optimization methods to obtain a point estimate for the model defined by S4 class `stanmodel` given the data, initial values, etc.

See Also

`stanmodel`

Examples

```r
## Not run:
m <- stan_model(model_code = 'parameters {real y;} model {y ~ normal(0,1);}')
f <- optimizing(m, hessian = TRUE)
## End(Not run)
```
Create a matrix of output plots from a `stanfit` object

Description

A `pairs` method that is customized for MCMC output

Usage

```r
## S3 method for class 'stanfit'
pairs(x, labels = NULL, panel = NULL, ..., 
lower.panel = NULL, 
upper.panel = NULL, diag.panel = NULL, text.panel = NULL, 
label.pos = 0.5 + 1/3, cex.labels = NULL, font.labels = 1, 
rowlattop = TRUE, gap = 1, log = "", pars = NULL, include = TRUE, 
condition = "accept_stat__")
```

Arguments

- `x` An object of S4 class `stanfit`
- `labels`, `panel`, `...`, `lower.panel`, `upper.panel`, `diag.panel` 
  Same as in `pairs` syntactically but see the Details section for different default arguments
- `text.panel`, `label.pos`, `cex.labels`, `font.labels`, `rowlattop`, `gap` 
  Same as in `pairs.default`
- `log` 
  Same as in `pairs.default`, which makes it possible to utilize logarithmic axes and additionally accepts `log = TRUE`. See the Details section.
- `pars` 
  If not NULL, a character vector indicating which quantities to include in the plots, which is passed to `extract`. Thus, by default, all unknown quantities are included, which may be far too many to visualize on a small computer screen. If `include = FALSE`, then the named parameters are excluded from the plot.
- `condition` 
  If NULL, it will plot roughly half of the chains in the lower panel and the rest in the upper panel. An integer vector can be passed to select some subset of the chains, of which roughly half will be plotted in the lower panel and the rest in the upper panel. A list of two integer vectors can be passed, each specifying a subset of the chains to be plotted in the lower and upper panels respectively. 
  A single number between zero and one exclusive can be passed, which is interpreted as the proportion of realizations (among all chains) to plot in the lower panel starting with the first realization in each chain, with the complement (from the end of each chain) plotted in the upper panel.
  A (possibly abbreviated) character vector of length one can be passed among "accept_stat__", "stepsize__", "treedepth__", "n_leapfrog__", "divergent__", "energy__", or "lp__", which are the variables produced by `get_sampler_params` and `get_logposterior`. In that case the lower panel will plot realizations that are below the median of the indicated variable (or are zero in the case
of "divergent__" and the upper panel will plot realizations that are greater than or equal to the median of the indicated variable (or are one in the case of "divergent__"). Finally, any logical vector whose length is equal to the product of the number of iterations and the number of chains can be passed, in which case realizations corresponding to FALSE and TRUE will be plotted in the lower and upper panel respectively. The default is "accept_stat__".

**include** Logical scalar indicating whether to include (the default) or exclude the parameters named in the `pars` argument from the plot.

**Details**

This method differs from the default `pairs` method in the following ways. If unspecified, the `smoothScatter` function is used for the off-diagonal plots, rather than `points`, since the former is more appropriate for visualizing thousands of draws from a posterior distribution. Also, if unspecified, histograms of the marginal distribution of each quantity are placed on the diagonal of the plot, after pooling all of the chains specified by the `chain_id` argument.

The draws from the warmup phase are always discarded before plotting.

By default, the lower (upper) triangle of the plot contains draws with below (above) median acceptance probability. Also, if condition is not "divergent__", red points will be superimposed onto the smoothed density plots indicating which (if any) iterations encountered a divergent transition. Otherwise, yellow points indicate a transition that hit the maximum treedepth rather than terminated its evolution normally.

You may very well want to specify the `log` argument for non-negative parameters. However, the `pairs` function will drop (with a message) parameters that are either constant or duplicative with previous parameters. For example, if a correlation matrix is included among `pars`, then neither its diagonal elements (which are always 1) nor its upper triangular elements (which are the same as the corresponding lower triangular elements) will be included. Thus, if `log` is an integer vector, it needs to pertain to the parameters after constant and duplicative ones are dropped. It is perhaps easiest to specify `log = TRUE`, which will utilize logarithmic axes for all non-negative parameters, except `lp__` and any integer valued quantities.

**See Also**

S4 class `stanfit` and its method `extract` as well as the `pairs` generic function. Also, see `get_sampler_params` and `get_logposterior`.

**Examples**

```r
example(read_stan_csv)
pairs(fit, pars = c("mu", "sigma", "alpha", "lp__"), log = TRUE, las = 1)
```

# sigma and alpha will have logarithmic axes
Description

The default plot shows posterior uncertainty intervals and point estimates for parameters and generated quantities. The plot method can also be used to call the other rstan plotting functions via the plotfun argument (see Examples).

Usage

```r
## S4 method for signature 'stanfit,missing'
plot(x, ..., plotfun)
```

Arguments

- `x` An instance of class `stanfit`.
- `plotfun` A character string naming the plotting function to apply to the stanfit object. If `plotfun` is missing, the default is to call `stan_plot`, which generates a plot of credible intervals and point estimates. See rstan-plotting-functions for the names and descriptions of the other plotting functions. `plotfun` can be either the full name of the plotting function (e.g. "stan_hist") or can be abbreviated to the part of the name following the underscore (e.g. "hist").
- `...` Optional arguments to `plotfun`.

Value

A `ggplot` object that can be further customized using the `ggplot2` package.

Note

Because the rstan plotting functions use `ggplot2` (and thus the resulting plots behave like ggplot objects), when calling a plotting function within a loop or when assigning a plot to a name (e.g., `graph <- plot(fit, plotfun = "rhat")`), if you also want the side effect of the plot being displayed you must explicitly print it (e.g., `(graph <- plot(fit, plotfun = "rhat")), print(graph <- plot(fit, plotfun = "rhat"))`).

See Also

List of RStan plotting functions, Plot options

Examples

```r
## Not run:
library(rstan)
fit <- stan_demo("eight_schools")
plot(fit)
plot(fit, show_density = TRUE, ci_level = 0.5, fill_color = "purple")
```
Plots

 ggplot2 for RStan

Description

Visual posterior analysis using ggplot2.

Usage

```r
stan_plot(object, pars, include = TRUE, unconstrain = FALSE, ...)
stan_trace(object, pars, include = TRUE, unconstrain = FALSE,
          inc_warmup = FALSE, nrow = NULL, ncol = NULL, ...,
          window = NULL)
stan_scat(object, pars, unconstrain = FALSE,
          inc_warmup = FALSE, nrow = NULL, ncol = NULL, ...)
stan_hist(object, pars, include = TRUE, unconstrain = FALSE,
          inc_warmup = FALSE, nrow = NULL, ncol = NULL, ...)
stan_dens(object, pars, include = TRUE, unconstrain = FALSE,
          inc_warmup = FALSE, nrow = NULL, ncol = NULL, ...)
stan_ac(object, pars, include = TRUE, unconstrain = FALSE,
          inc_warmup = FALSE, nrow = NULL, ncol = NULL, ..., 
          separate_chains = FALSE)
quietgg(gg)
```

Arguments

- `object` A stanfit or stanreg object.
- `pars` Optional character vector of parameter names. If `object` is a stanfit object, the default is to show all user-defined parameters or the first 10 (if there are more than 10). If `object` is a stanreg object, the default is to show all (or the first 10) regression coefficients (including the intercept). For `stan_scat` only, `pars` should not be missing and should contain exactly two parameter names.
- `include` Should the parameters given by the `pars` argument be included (the default) or excluded from the plot?
- `unconstrain` Should parameters be plotted on the unconstrained space? Defaults to FALSE. Only available if `object` is a stanfit object.
- `inc_warmup` Should warmup iterations be included? Defaults to FALSE.
- `nrow`, `ncol` Passed to `facet_wrap`.
Plots

... Optional additional named arguments passed to geoms (e.g. for stan_trace the geom is geom_path and we could specify linetype, size, alpha, etc.). For stan_plot there are also additional arguments that can be specified in ... (see Details).

window For stan_trace window is used to control which iterations are shown in the plot. See traceplot.

separate_chains For stan_dens, should the density for each chain be plotted? The default is FALSE, which means that for each parameter the draws from all chains are combined. For stan_ac, if separate_chains=FALSE (the default), the autocorrelation is averaged over the chains. If TRUE each chain is plotted separately.

lags For stan_ac, the maximum number of lags to show.

partial For stan_ac, should partial autocorrelations be plotted instead? Defaults to FALSE.

gg A ggplot object or an expression that creates one.

Details

For stan_plot, there are additional arguments that can be specified in ... The optional arguments and their default values are:

point_est = "median" The point estimate to show. Either "median" or "mean".

show_density = FALSE Should kernel density estimates be plotted above the intervals?

ci_level = 0.8 The posterior uncertainty interval to highlight. Central 100*ci_level% intervals are computed from the quantiles of the posterior draws.

outer_level = 0.95 An outer interval to also draw as a line (if show_outer_line is TRUE) but not highlight.

show_outer_line = TRUE Should the outer_level interval be shown or hidden? Defaults to = TRUE (to plot it).

fill_color, outline_color, est_color Colors to override the defaults for the highlighted interval, the outer interval (and density outline), and the point estimate.

Value

A ggplot object that can be further customized using the ggplot2 package.

Note

Because the rstan plotting functions use ggplot2 (and thus the resulting plots behave like ggplot objects), when calling a plotting function within a loop or when assigning a plot to a name (e.g., graph <- plot(fit, plotfun = "rhat")), if you also want the side effect of the plot being displayed you must explicitly print it (e.g., (graph <- plot(fit, plotfun = "rhat")), print(graph <- plot(fit, plotfun = "rhat"))).

See Also

List of RStan plotting functions, Plot options
Examples

```r
## Not run:
example("read_stan_csv")
stan_plot(fit)
stan_trace(fit)

library(gridExtra)
fit <- stan_demo("eight_schools")

stan_plot(fit)
stan_plot(fit, point_est = "mean", show_density = TRUE, fill_color = "maroon")

# histograms
stan_hist(fit)
# suppress ggplot2 messages about default bindwidth
quietgg(stan_hist(fit))
quietgg(h <- stan_hist(fit, pars = "theta", binwidth = 5))

# juxtapose histograms of tau and unconstrained tau
tau <- stan_hist(fit, pars = "tau")
tau_unc <- stan_hist(fit, pars = "tau", unconstrain = TRUE) +
  xlab("tau unconstrained")
grid.arrange(tau, tau_unc)

# kernel density estimates
stan_dens(fit)
(dens <- stan_dens(fit, fill = "skyblue", ))
dens <- dens + ggtitle("Kernel Density Estimates\n") + xlab("")
dens

(dens_sep <- stan_dens(fit, separate_chains = TRUE, alpha = 0.3))
dens_sep + scale_fill_manual(values = c("red", "blue", "green", "black"))
(dens_sep_stack <- stan_dens(fit, pars = "theta", alpha = 0.5,
  separate_chains = TRUE, position = "stack"))

# traceplot
trace <- stan_trace(fit)
trace +
  scale_color_manual(values = c("red", "blue", "green", "black"))
trace +
  scale_color_brewer(type = "div") +
  theme(legend.position = "none")

facet_style <- theme(strip.background = ggplot2::element_rect(fill = "white"),
  strip.text = ggplot2::element_text(size = 13, color = "black"))
(trace <- trace + facet_style)

# scatterplot
(mu_vs_tau <- stan_scat(fit, pars = c("mu", "tau"), color = "blue", size = 4))
mu_vs_tau +
  ggplot2::coord_flip()
```
print

theme(panel.background = ggplot2::element_rect(fill = "black"))

## End(Not run)

---

**print**  
*Print a summary for a fitted model represented by a stanfit object*

---

**Description**

Print basic information regarding the fitted model and a summary for the parameters of interest estimated by the samples included in a `stanfit` object.

**Usage**

```r
## S3 method for class 'stanfit'
print(x, pars = x@sim$pars_oi,
       probs = c(0.025, 0.25, 0.5, 0.75, 0.975),
       digits_summary = 2, include = TRUE, ...)  
```

**Arguments**

- `x`  
  An object of S4 class `stanfit`.

- `pars`  
  A character vector of parameter names. The default is all parameters for which samples are saved. If `include = FALSE`, then the specified parameters are excluded from the printed summary.

- `probs`  
  A numeric vector of quantiles of interest. The default is `c(0.025, 0.25, 0.5, 0.75, 0.975)`.  

- `digits_summary`  
  The number of significant digits to use when printing the summary, defaulting to 2. Applies to the quantities other than the effective sample size, which is always rounded to the nearest integer.

- `include`  
  Logical scalar (defaulting to `TRUE`) indicating whether to include or exclude the parameters named by the `pars` argument.

- `...`  
  Additional arguments passed to the `summary` method for `stanfit` objects.

**Details**

The information regarding the fitted model includes the number of iterations, the number of chains, the total number of saved iterations, the estimation algorithm used, and the timestamp indicating when sampling finished.

The parameter summaries computed include means, standard deviations (sd), quantiles, Monte Carlo standard errors (se_mean), split Rhats, and effective sample sizes (n_eff). The summaries are computed after dropping the warmup iterations and merging together the draws from all chains.

In addition to the model parameters, summaries for the log-posterior (lp__) are also reported.
read_rdump

See Also

S4 class stanfit and particularly its method summary, which is used to obtain the values that are printed.

read_rdump

Read data in an R dump file to a list

Description

Create an R list from an R dump file

Usage

read_rdump(f, keep.source = FALSE, ...)

Arguments

  f
    A character string providing the dump file name.

  keep.source
    logical: should the source formatting be retained when echoing expressions, if possible?

  ...
    passed to source

Details

The R dump file can be read directly by R function source, which by default would read the data into the user's workspace (the global environment). This function instead read the data to a list, making it convenient to prepare data for the stan model-fitting function.

Value

A list containing all the data defined in the dump file with keys corresponding to variable names.

See Also

stan_rdump; dump

Examples

x <- 1; y <- 1:10; z <- array(1:10, dim = c(2,5))
stan_rdump(ls(pattern = "^[xyz]"), file.path(tempdir(), "xyz.Rdump"))
l <- read_rdump(file.path(tempdir(), "xyz.Rdump"))
print(l)
unlink(file.path(tempdir(), "xyz.Rdump"))
read_stan_csv

Read CSV files of samples generated by (R)Stan into a stanfit object

Description

Create a stanfit object from the saved CSV files that are created by Stan or RStan and that include the samples drawn from the distribution of interest to facilitate analysis of samples using RStan.

Usage

read_stan_csv(csvfiles, col_major = TRUE)

Arguments

csvfiles: A character vector providing CSV file names

col_major: The order for array parameters; default to TRUE

Details

Stan and RStan could save the samples to CSV files. This function reads the samples and using the comments (beginning with "#") to create a stanfit object. The model name is derived from the first CSV file.

col_major specifies how array parameters are ordered in each row of the CSV files. For example, parameter "a[2,2]" would be ordered as "a[1,1], a[2,1], a[1,2], a[2,2]" if col_major is TRUE.

Value

A stanfit object (with invalid stanmodel slot). This stanfit object cannot be used to re-run the sampler.

See Also

stanfit

Examples

csvfiles <- dir(system.file('misc', package = 'rstan'), pattern = 'rstan_doc_ex_[0-9].csv', full.names = TRUE)
fit <- read_stan_csv(csvfiles)
**Rhat**

*Convergence and efficiency diagnostics for Markov Chains*

---

**Description**

These functions are improved versions of the traditional Rhat (for convergence) and Effective Sample Size (for efficiency).

**Usage**

```r
Rhat(sims)
ess_bulk(sims)
ess_tail(sims)
```

**Arguments**

- `sims`: A two-dimensional array whose rows are equal to the number of iterations of the Markov Chain(s) and whose columns are equal to the number of Markov Chains (preferably more than one). The cells are the realized draws for a particular parameter or function of parameters.

**Value**

The `Rhat` function produces R-hat convergence diagnostic, which compares the between- and within-chain estimates for model parameters and other univariate quantities of interest. If chains have not mixed well (i.e., the between- and within-chain estimates don’t agree), R-hat is larger than 1. We recommend running at least four chains by default and only using the sample if R-hat is less than 1.05. Stan reports R-hat which is the maximum of rank normalized split-R-hat and rank normalized folded-split-R-hat, which works for thick tailed distributions and is sensitive also to differences in scale.

The `ess_bulk` function produces an estimated Bulk Effective Sample Size (bulk-ESS) using rank normalized draws. Bulk-ESS is useful measure for sampling efficiency in the bulk of the distribution (related e.g., to efficiency of mean and median estimates), and is well defined even if the chains do not have finite mean or variance.

The `ess_tail` function produces an estimated Tail Effective Sample Size (tail-ESS) by computing the minimum of effective sample sizes for 5% and 95% quantiles. Tail-ESS is useful measure for sampling efficiency in the tails of the distribution (related e.g., to efficiency of variance and tail quantile estimates).

Both bulk-ESS and tail-ESS should be at least 100 (approximately) per Markov Chain in order to be reliable and indicate that estimates of respective posterior quantiles are reliable.

**Author(s)**

Paul-Christian Burkner and Aki Vehtari
References


See Also

`monitor`

Examples

```r
# pretend these draws came from five actual Markov Chins
sims <- matrix(rnorm(500), nrow = 100, ncol = 5)
Rhat(sims)
ess_bulk(sims)
ess_tail(sims)
```

---

**rstan-plotting-functions**

*RStan Plotting Functions*

---

Description

List of RStan plotting functions that return ggplot objects

**RStan plotting functions**

- **Posterior intervals and point estimates** `stan_plot`
- **Traceplots** `stan_trace`
- **Histograms** `stan_hist`
- **Kernel density estimates** `stan_dens`
- **Scatterplots** `stan_scat`
- **Diagnostics for Hamiltonian Monte Carlo and the No-U-Turn Sampler** `stan_diag`
- **Rhat** `stan_rhat`
- **Ratio of effective sample size to total posterior sample size** `stan_ess`
- **Ratio of Monte Carlo standard error to posterior standard deviation** `stan_mcse`
- **Autocorrelation** `stan_ac`

See Also

`Plot options`
rstan.package.skeleton

Create a Skeleton for a New Source Package with Stan Programs

Description

This function has been removed from rstan. Please use the new rstan_package_skeleton function in the rstantools package.

rstan_gg_options

Set default appearance options

Description

Set default appearance options

Usage

rstan_gg_options(...)

rstan_ggtheme_options(...)

Arguments

... For rstan_ggtheme_options, see theme for the theme elements that can be specified in .... For rstan_gg_options, ... can be fill.color, chain_colors, size, pt_color, or pt_size. See Examples.

See Also

List of RStan plotting functions

Examples

rstan_ggtheme_options(panel.background = ggplot2::element_rect(fill = "gray"),
                       legend.position = "top")
rstan_gg_options(fill = "skyblue", color = "skyblue4", pt_color = "red")
rstan_options

Set and read options used in RStan

Description
Set and read options used in RStan. Some settings as options can be controlled by the user.

Usage
rstan_options(...)

Arguments

... Arguments of the form opt = val set option opt to value val. Arguments of
the form opt set the function to return option opt's value. Each argument must
be a character string.

Details
The available options are:

1. plot_rhat_breaks: The cut off points for Rhat for which we would indicate using a different
   color. This is a numeric vector, defaulting to c(1.1, 1.2, 1.5, 2). The value for this
   option will be sorted in ascending order, so for example plot_rhat_breaks = c(1.2, 1.5)
   is equivalent to plot_rhat_breaks = c(1.5, 1.2).
2. plot_rhat_cols: A vector of the same length as plot_rhat_breaks that indicates the colors
   for the breaks.
3. plot_rhat_nan_col: The color for Rhat when it is Inf or NaN.
4. plot_rhat_large_col: The color for Rhat when it is larger than the largest value of plot_rhat_breaks.
5. rstan_alert_col: The color used in method plot of S4 class stanfit to show that the
   vector/array parameters are truncated.
6. rstan_chain_cols: The colors used in methods plot and traceplot of S4 class stanfit
   for coloring different chains.
7. rstan_warmup_bg_col: The background color for the warmup area in the traceplots.
8. boost_lib: The path for the Boost C++ library used to compile Stan models. This option is
   valid for the whole R session if not changed again.
9. eigen_lib: The path for the Eigen C++ library used to compile Stan models. This option is
   valid for the whole R session if not changed again.
10. auto_write: A logical scalar (defaulting to FALSE) that controls whether a compiled instance
   of a stanmodel-class is written to the hard disk in the same directory as the .stan program.

Value
The values as a list for existing options and NA for non-existent options. When only one option is
specified, its old value is returned.
**Description**

Draw samples from the model defined by class stanmodel.

**Usage**

```r
## S4 method for signature 'stanmodel'
sampling(object, data = list(), pars = NA,
  chains = 4, iter = 2000, warmup = floor(iter/2), thin = 1,
  seed = sample.int(.Machine$integer.max, 1),
  init = 'random', check.data = TRUE,
  sample_file = NULL, diagnostic_file = NULL, verbose = FALSE,
  algorithm = c("NUTS", "HMC", "Fixed_param"),
  control = NULL, include = TRUE,
  cores = getOption("mc.cores", 1L),
  open_progress = interactive() && !isatty(stdout()) &&
    !identical(Sys.getenv("RSTUDIO"), "1"),
  show_messages = TRUE, ...)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>An object of class <code>stanmodel</code>.</td>
</tr>
<tr>
<td>data</td>
<td>A named list or environment providing the data for the model or a character vector for all the names of objects used as data. See the Passing data to Stan section in <code>stan</code>.</td>
</tr>
<tr>
<td>pars</td>
<td>A vector of character strings specifying parameters of interest. The default is <code>NA</code> indicating all parameters in the model. If <code>include = TRUE</code>, only samples for parameters named in <code>pars</code> are stored in the fitted results. Conversely, if <code>include = FALSE</code>, samples for all parameters except those named in <code>pars</code> are stored in the fitted results.</td>
</tr>
<tr>
<td>chains</td>
<td>A positive integer specifying the number of Markov chains. The default is 4.</td>
</tr>
<tr>
<td>iter</td>
<td>A positive integer specifying the number of iterations for each chain (including warmup). The default is 2000.</td>
</tr>
<tr>
<td>warmup</td>
<td>A positive integer specifying the number of warmup (aka burnin) iterations per chain. If step-size adaptation is on (which it is by default), this also controls the number of iterations for which adaptation is run (and hence these warmup samples should not be used for inference). The number of warmup iterations should not be larger than <code>iter</code> and the default is <code>iter/2</code>.</td>
</tr>
<tr>
<td>thin</td>
<td>A positive integer specifying the period for saving samples. The default is 1, which is usually the recommended value.</td>
</tr>
</tbody>
</table>
seed  The seed for random number generation. The default is generated from 1 to the
maximum integer supported by R on the machine. Even if multiple chains are
used, only one seed is needed, with other chains having seeds derived from that
of the first chain to avoid dependent samples. When a seed is specified by a
number, as.integer will be applied to it. If as.integer produces NA, the seed
is generated randomly. The seed can also be specified as a character string of
digits, such as "12345", which is converted to integer.

init  Initial values specification. See the detailed documentation for the init argument
in stan.

check_data  Logical, defaulting to TRUE. If TRUE the data will be preprocessed; otherwise
not. See the Passing data to Stan section in stan.

time_file  An optional character string providing the name of a file. If specified the draws
for all parameters and other saved quantities will be written to the file. If not pro-
vided, files are not created. When the folder specified is not writable, tempdir() is
used. When there are multiple chains, an underscore and chain number are
appended to the file name prior to the .csv extension.

diagnostic_file  An optional character string providing the name of a file. If specified the diag-
nostics data for all parameters will be written to the file. If not provided, files
are not created. When the folder specified is not writable, tempdir() is used.
When there are multiple chains, an underscore and chain number are appended
to the file name prior to the .csv extension.

verbose  TRUE or FALSE: flag indicating whether to print intermediate output from Stan
on the console, which might be helpful for model debugging.

algorithm  One of sampling algorithms that are implemented in Stan. Current options are
"NUTS" (No-U-Turn sampler, Hoffman and Gelman 2011, Betancourt 2017),
"HMC" (static HMC), or "fixed_param". The default and preferred algorithm is
"NUTS".

control  A named list of parameters to control the sampler’s behavior. See the details
in the documentation for the control argument in stan.

include  Logical scalar defaulting to TRUE indicating whether to include or exclude the
parameters given by the pars argument. If FALSE, only entire multidimensional
parameters can be excluded, rather than particular elements of them.

cores  Number of cores to use when executing the chains in parallel, which defaults to
1 but we recommend setting the mc.cores option to be as many processors as
the hardware and RAM allow (up to the number of chains).

open_progress  Logical scalar that only takes effect if cores > 1 but is recommended to be
TRUE in interactive use so that the progress of the chains will be redirected to
a file that is automatically opened for inspection. For very short runs, the user
might prefer FALSE.

show_messages  Either a logical scalar (defaulting to TRUE) indicating whether to print the sum-
mary of Informational Messages to the screen after a chain is finished or a char-
acter string naming a path where the summary is stored. Setting to FALSE is not
recommended unless you are very sure that the model is correct up to numerical
error.
Additional arguments can be chain_id, init_r, test_grad, append_samples, refresh, enable_random_init. See the documentation in `stan`.

Value

An object of S4 class `stanfit` representing the fitted results. Slot `mode` for this object indicates if the sampling is done or not.

Methods

`sampling` signature(`object = "stanmodel"`)

Call a sampler (NUTS, HMC, or Fixed_param depending on parameters) to draw samples from the model defined by S4 class `stanmodel` given the data, initial values, etc.

See Also

`stanmodel`, `stanfit`, `stan`

Examples

```r
## Not run:
m <- stan_model(model_code = 'parameters {real y;} model {y ~ normal(0,1);}')
f <- sampling(m, iter = 100)
## End(Not run)
```

---

**sbc**

*Simulation Based Calibration (sbc)*

Description

Check whether a model is well-calibrated with respect to the prior distribution and hence possibly amenable to obtaining a posterior distribution conditional on observed data.

Usage

```r
sbc(stanmodel, data, M, ...)
## S3 method for class 'sbc'
plot(x, thin = 3, ...)
## S3 method for class 'sbc'
print(x, ...)
```
Arguments

stanmodel  An object of stanmodel-class that is first created by calling the stan_model function

data  A named list or environment providing the data for the model, or a character vector for all the names of objects to use as data. This is the same format as in stan or sampling.

M  The number of times to condition on draws from the prior predictive distribution

...  Additional arguments that are passed to sampling, such as refresh = 0 when calling sbc. For the plot and print methods, the additional arguments are not used.

x  An object produced by sbc

thin  An integer vector of length one indicating the thinning interval when plotting, which defaults to 3

Details

This function assumes adherence to the following conventions in the underlying Stan program:

1. Realizations of the unknown parameters are drawn in the transformed data block of the Stan program and are postfixed with an underscore, such as theta_. These are considered the “true” parameters being estimated by the corresponding symbol declared in the parameters block, which should have the same name except for the trailing underscore, such as theta.

2. The realizations of the unknown parameters are then conditioned on when drawing from the prior predictive distribution, also in the transformed data block. There is no restriction on the symbol name that holds the realizations from the prior predictive distribution but for clarity, it should not end with a trailing underscore.

3. The realizations of the unknown parameters should be copied into a vector in the generated quantities block named pars_.

4. The realizations from the prior predictive distribution should be copied into an object (of the same type) in the generated quantities block named y_. Technically, this step is optional and could be omitted to conserve RAM, but inspecting the realizations from the prior predictive distribution is a good way to judge whether the priors are reasonable.

5. The generated quantities block must contain an integer array named ranks_ whose only values are zero or one, depending on whether the realization of a parameter from the posterior distribution exceeds the corresponding “true” realization, such as theta > theta_. These are not actually "ranks" but can be used afterwards to reconstruct (thinned) ranks.

6. The generated quantities block may contain a vector named log_lik whose values are the contribution to the log-likelihood by each observation. This is optional but facilitates calculating Pareto k shape parameters to judge whether the posterior distribution is sensitive to particular observations.

Although the user can pass additional arguments to sampling through the ..., the following arguments are hard-coded and should not be passed through the ...:

1. pars = "ranks_" because nothing else needs to be stored for each posterior draw

2. include = TRUE to ensure that "ranks_" is included rather than excluded
3. chains = 1 because only one chain is run for each integer less than M
4. seed because a sequence of seeds is used across the M runs to preserve independence across runs
5. save_warmup = FALSE because the warmup realizations are not relevant
6. thin = 1 because thinning can and should be done after the Markov Chain is finished, as is done by the thin argument to the plot method in order to make the histograms consist of approximately independent realizations

Other arguments will take the default values used by sampling unless passed through the .... Specifying refresh = 0 is recommended to avoid printing a lot of intermediate progress reports to the screen. It may be necessary to pass a list to the control argument of sampling with elements adapt_delta and / or max_treedepth in order to obtain adequate results.

Ideally, users would want to see the absence of divergent transitions (which is shown by the print method) and other warnings, plus an approximately uniform histogram of the ranks for each parameter (which are shown by the plot method). See the vignette for more details.

Value

The sbc function outputs a list of S3 class "sbc", which contains the following elements:

1. ranks A list of M matrices, each with number of rows equal to the number of saved iterations and number of columns equal to the number of unknown parameters. These matrices contain the realizations of the ranks_ object from the generated quantities block of the Stan program.
2. y If present, a matrix of realizations from the prior predictive distribution whose rows are equal to the number of observations and whose columns are equal to M, which are taken from the y_ object in the generated quantities block of the Stan program.
3. pars A matrix of realizations from the prior distribution whose rows are equal to the number of parameters and whose columns are equal to M, which are taken from the pars_ object in the generated quantities block of the Stan program.
4. pareto_k A matrix of Pareto k shape parameter estimates or NULL if there is no log_lik symbol in the generated quantities block of the Stan program.
5. sampler_params A three-dimensional array that results from combining calls to get_sampler_params for each of the M runs. The resulting matrix has rows equal to the number of post-warmup iterations, columns equal to six, and M floors. The columns are named "accept_stat__", "stepsize__", "treedepth__", "n_leapfrog__", "divergent__", and "energy__". The most important of which is "divergent__", which should be all zeros and perhaps "treedepth__", which should only rarely get up to the value of max_treedepth passed as an element of the control list to sampling or otherwise defaults to 10.

The print method outputs the number of divergent transitions and returns NULL invisibly. The plot method returns a ggplot object with histograms whose appearance can be further customized.

References


See Also

`stan_model` and `sampling`

Examples

```r
scode <- "
data {
  int<lower = 1> N;
  real<lower = 0> a;
  real<lower = 0> b;
}
transformed data { // these adhere to the conventions above
  real pi_ = beta_rng(a, b);
  int y = binomial_rng(N, pi_);
}
parameters {
  real<lower = 0, upper = 1> pi;
}
model {
  target += beta_lpdf(pi | a, b);
  target += binomial_lpmf(y | N, pi);
}
generated quantities { // these adhere to the conventions above
  int y_ = y;
  vector[1] pars_;
  int ranks_[1] = (pi > pi_);
  vector[N] log_lik;
  pars_[1] = pi_;
  for (n in 1:y) log_lik[n] = bernoulli_lpmf(1 | pi);
  for (n in (y + 1):N) log_lik[n] = bernoulli_lpmf(0 | pi);
}"
```

---

**set_cppo**  
*Defunct function to set the compiler optimization level*

**Description**

This function returns nothing and does nothing except throw a warning. See https://cran.r-project.org/doc/manuals/r-release/R-admin.html#Customizing-package-compilation for information on customizing the compiler options, but doing so should be unnecessary for normal usage.
sflist2stanfit

Usage

```r
set_cppo(...)```

Arguments

```r
... Any input is ignored```

Value

An invisible NULL

---

`sflist2stanfit` *Merge a list of stanfit objects into one*

Description

This function takes a list of `stanfit` objects and returns a consolidated `stanfit` object. The `stanfit` objects to be merged need to have the same configuration of iteration, warmup, and thin, besides being from the same model. This could facilitate some parallel usage of RStan. For example, if we call `stan` by parallel and it returns a list of `stanfit` objects, this function can be used to create one `stanfit` object from the list.

Usage

```r
sflist2stanfit(sflist)
```

Arguments

```r
sflist A list of stanfit objects.
```

Value

An S4 object of `stanfit` consolidated from all the input `stanfit` objects.

Note

This function should be called in rare circumstances because `sampling` has a `cores` argument that allows multiple chains to be executed in parallel. However, if you need to depart from that, the best practice is to use `sflist2stanfit` on a list of `stanfit` objects created with the same seed but different `chain_id` (see example below). Using the same seed but different `chain_id` can make sure the random number generations for all chains are not correlated.

This function would do some check to see if the `stanfit` objects in the input list can be merged. But the check is not sufficient. So generally, it is the user’s responsibility to make sure the input is correct so that the merging makes sense.

The date in the new `stanfit` object is when it is merged.

`get_seed` function for the new consolidated `stanfit` object only returns the seed used in the first chain of the new object.
The sampler such as NUTS2 that is displayed in the printout by `print` is the sampler used for the first chain. The `print` method assumes the samplers are the same for all chains.

The included `stanmodel` object, which includes the compiled model, in the new `stanfit` object is from the first element of the input list.

References


See Also

`stan`

Examples

```r
## Not run:
library(rstan)
scode <- "
data {
  int<lower=1> N;
}
parameters {
  real y1[N];
  real y2[N];
}
model {
  y1 ~ normal(0, 1);
  y2 ~ double_exponential(0, 2);
}
"
seed <- 123 # or any other integer
foo_data <- list(N = 2)
foo <- stan(model_code = scode, data = foo_data, chains = 1, iter = 1)
f1 <- stan(fit = foo, data = foo_data, chains = 1, seed = seed, chain_id = 1)
f2 <- stan(fit = foo, data = foo_data, chains = 2, seed = seed, chain_id = 2:3)
f12 <- sflist2stanfit(list(f1, f2))

## parallel stan call for unix-like OS
library(parallel)

if (.Platform$OS.type == "unix") {
  sflist1 <-
    mclapply(1:4, mc.cores = 2,
              function(i) stan(fit = foo, data = foo_data, seed = seed,
                               chains = 1, chain_id = i, refresh = -1))
  f3 <- sflist2stanfit(sflist1)
}
if (.Platform$OS.type == "windows") { # also works on non-Windows
  CL <- makeCluster(2)
  clusterExport(CL = CL, c("foo_data", "foo", "seed"))
  sflist1 <- parLapply(CL, 1:4, fun = function(cid) {
```
require(rstan)
stan(fit = foo, data = foo_data, chains = 1,
    iter = 2000, seed = seed, chain_id = cid)
}

fit <- sflist2stanfit(sflist)
print(fit)
stopCluster(CL)
} # end example for Windows

## End(Not run)

---

### Stan

Fit a model with Stan

#### Description

Fit a model defined in the Stan modeling language and return the fitted result as an instance of stanfit.

#### Usage

```r
stan(file, model_name = "anon_model", model_code = "", fit = NA,
    data = list(), pars = NA,
    chains = 4, iter = 2000, warmup = floor(iter/2), thin = 1,
    init = "random", seed = sample.int(.Machine$integer.max, 1),
    algorithm = c("NUTS", "HMC", "Fixed_param"),
    control = NULL, sample_file = NULL, diagnostic_file = NULL,
    save_dso = TRUE, verbose = FALSE, include = TRUE,
    cores = getOption("mc.cores", 1L),
    open_progress = interactive() && !isatty(stdout()) &&
        !identical(Sys.getenv("RSTUDIO"), "1"),
    ...
    boost_lib = NULL, eigen_lib = NULL
)
```

#### Arguments

- **file**: The path to the Stan program to use. file should be a character string file name or a connection that R supports containing the text of a model specification in the Stan modeling language.

  A model may also be specified directly as a character string using the model_code argument, but we recommend always putting Stan programs in separate files with a .stan extension.

  The stan function can also use the Stan program from an existing stanfit object via the fit argument. When fit is specified, the file argument is ignored.
<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>model_code</code></td>
<td>A character string either containing the model definition or the name of a character string object in the workspace. This argument is used only if arguments <code>file</code> and <code>fit</code> are not specified.</td>
</tr>
<tr>
<td><code>fit</code></td>
<td>An instance of S4 class <code>stanfit</code> derived from a previous fit; defaults to <code>NA</code>. If <code>fit</code> is not <code>NA</code>, the compiled model associated with the fitted result is re-used; thus the time that would otherwise be spent recompiling the C++ code for the model can be saved.</td>
</tr>
<tr>
<td><code>model_name</code></td>
<td>A string to use as the name of the model; defaults to &quot;anon_model&quot;. However, the model name will be derived from <code>file</code> or <code>model_code</code> (if <code>model_code</code> is the name of a character string object) if <code>model_name</code> is not specified. This is not a particularly important argument, although since it affects the name used in printed messages, developers of other packages that use rstan to fit models may want to use informative names.</td>
</tr>
<tr>
<td><code>data</code></td>
<td>A named list or environment providing the data for the model, or a character vector for all the names of objects to use as data. See the <strong>Passing data to Stan</strong> section below.</td>
</tr>
<tr>
<td><code>pars</code></td>
<td>A character vector specifying parameters of interest to be saved. The default is to save all parameters from the model. If <code>include = TRUE</code>, only samples for parameters named in <code>pars</code> are stored in the fitted results. Conversely, if <code>include = FALSE</code>, samples for all parameters except those named in <code>pars</code> are stored in the fitted results.</td>
</tr>
<tr>
<td><code>include</code></td>
<td>Logical scalar defaulting to <code>TRUE</code> indicating whether to include or exclude the parameters given by the <code>pars</code> argument. If <code>FALSE</code>, only entire multidimensional parameters can be excluded, rather than particular elements of them.</td>
</tr>
<tr>
<td><code>iter</code></td>
<td>A positive integer specifying the number of iterations for each chain (including warmup). The default is 2000.</td>
</tr>
<tr>
<td><code>warmup</code></td>
<td>A positive integer specifying the number of warmup (aka burnin) iterations per chain. If step-size adaptation is on (which it is by default), this also controls the number of iterations for which adaptation is run (and hence these warmup samples should not be used for inference). The number of warmup iterations should not be larger than <code>iter</code> and the default is <code>iter/2</code>.</td>
</tr>
<tr>
<td><code>chains</code></td>
<td>A positive integer specifying the number of Markov chains. The default is 4.</td>
</tr>
<tr>
<td><code>cores</code></td>
<td>The number of cores to use when executing the Markov chains in parallel. The default is to use the value of the &quot;mc.cores&quot; option if it has been set and otherwise to default to 1 core. However, we recommend setting it to be as many processors as the hardware and RAM allow (up to the number of chains). See <code>detectCores</code> if you don’t know this number for your system.</td>
</tr>
<tr>
<td><code>thin</code></td>
<td>A positive integer specifying the period for saving samples. The default is 1, which is usually the recommended value. Unless your posterior distribution takes up too much memory we do not recommend thinning as it throws away information. The tradition of thinning when running MCMC stems primarily from the use of samplers that require a large number of iterations to achieve the desired effective sample size. Because of the efficiency (effective samples per second) of Hamiltonian Monte Carlo, rarely should this be necessary when using Stan.</td>
</tr>
</tbody>
</table>
Specification of initial values for all or some parameters. Can be the digit 0, the strings "0" or "random", a function that returns a named list, or a list of named lists:

- `init="random"` (default): Let Stan generate random initial values for all parameters. The seed of the random number generator used by Stan can be specified via the `seed` argument. If the seed for Stan is fixed, the same initial values are used. The default is to randomly generate initial values between $-2$ and $2$ on the unconstrained support. The optional additional parameter `init_r` can be set to some value other than 2 to change the range of the randomly generated inits.
- `init="0", init=0`: Initialize all parameters to zero on the unconstrained support.

- `inits via list`: Set initial values by providing a list equal in length to the number of chains. The elements of this list should themselves be named lists, where each of these named lists has the name of a parameter and is used to specify the initial values for that parameter for the corresponding chain.

- `inits via function`: Set initial values by providing a function that returns a list for specifying the initial values of parameters for a chain. The function can take an optional parameter `chain_id` through which the `chain_id` (if specified) or the integers from 1 to `chains` will be supplied to the function for generating initial values. See the Examples section below for examples of defining such functions and using a list of lists for specifying initial values.

When specifying initial values via a list or function, any parameters for which values are not specified will receive initial values generated as described in the `init="random"` description above.

The seed for random number generation. The default is generated from 1 to the maximum integer supported by R on the machine. Even if multiple chains are used, only one seed is needed, with other chains having seeds derived from that of the first chain to avoid dependent samples. When a seed is specified by a number, `as.integer` will be applied to it. If `as.integer` produces `NA`, the seed is generated randomly. The seed can also be specified as a character string of digits, such as "12345", which is converted to integer.

Using R’s `set.seed` function to set the seed for Stan will not work.

One of the sampling algorithms that are implemented in Stan. The default and preferred algorithm is "NUTS", which is the No-U-Turn sampler variant of Hamiltonian Monte Carlo (Hoffman and Gelman 2011, Betancourt 2017). Currently the other options are "HMC" (Hamiltonian Monte Carlo), and "Fixed_param". When "Fixed_param" is used no MCMC sampling is performed (e.g., for simulating with in the generated quantities block).

An optional character string providing the name of a file. If specified the draws for all parameters and other saved quantities will be written to the file. If not provided, files are not created. When the folder specified is not writable, `tempdir()` is used. When there are multiple chains, an underscore and chain number are appended to the file name.

An optional character string providing the name of a file. If specified the diagnostics data for all parameters will be written to the file. If not provided, files
are not created. When the folder specified is not writable, `tempdir()` is used.
When there are multiple chains, an underscore and chain number are appended
to the file name.

**save_dso**
Logical, with default `TRUE`, indicating whether the dynamic shared object (DSO)
compiled from the C++ code for the model will be saved or not. If `TRUE`, we can
draw samples from the same model in another R session using the saved DSO
(i.e., without compiling the C++ code again). This parameter only takes effect if
`fit` is not used; with `fit` defined, the DSO from the previous run is used. When
`save_dso=TRUE`, the fitted object can be loaded from what is saved previously
and used for sampling, if the compiling is done on the same platform, that is,
same operating system and same architecture (32bits or 64bits).

**verbose**
`TRUE` or `FALSE`: flag indicating whether to print intermediate output from Stan
on the console, which might be helpful for model debugging.

**control**
A named list of parameters to control the sampler’s behavior. It defaults to
`NULL` so all the default values are used. First, the following are adaptation parameters
for sampling algorithms. These are parameters used in Stan with similar
names here.

- `adapt_engaged` (logical)
- `adapt_gamma` (double, positive, defaults to 0.05)
- `adapt_delta` (double, between 0 and 1, defaults to 0.8)
- `adapt_kappa` (double, positive, defaults to 0.75)
- `adapt_t0` (double, positive, defaults to 10)
- `adapt_init_buffer` (integer, positive, defaults to 75)
- `adapt_term_buffer` (integer, positive, defaults to 50)
- `adapt_window` (integer, positive, defaults to 25)

In addition, algorithm HMC (called 'static HMC' in Stan) and NUTS share the
following parameters:

- `stepsize` (double, positive)
- `stepsize_jitter` (double, [0,1])
- `metric` (string, one of "unit_e", "diag_e", "dense_e")

For algorithm NUTS, we can also set:

- `max_treedepth` (integer, positive)

For algorithm HMC, we can also set:

- `int_time` (double, positive)

For test_grad mode, the following parameters can be set:

- `epsilon` (double, defaults to 1e-6)
- `error` (double, defaults to 1e-6)

**open_progress**
Logical scalar that only takes effect if `cores > 1` but is recommended to be
`TRUE` in interactive use so that the progress of the chains will be redirected to
a file that is automatically opened for inspection. For very short runs, the user
might prefer `FALSE`.

... Other optional parameters:
chain_id (integer)
init_r (double, positive)
test_grad (logical)
append_samples (logical)
refresh (integer)
save_warmup (logical)
deprecated: enable_random_init (logical)

chain_id can be a vector to specify the chain_id for all chains or an integer. For the former case, they should be unique. For the latter, the sequence of integers starting from the given chain_id are used for all chains.

init_r is used only for generating random initial values, specifically when init="random" or not all parameters are initialized in the user-supplied list or function. If specified, the initial values are simulated uniformly from interval [-init_r, init_r] rather than using the default interval (see the manual of (cmd)Stan).

test_grad (logical). If test_grad=TRUE, Stan will not do any sampling. Instead, the gradient calculation is tested and printed out and the fitted stanfit object is in test gradient mode. By default, it is FALSE.

append_samples (logical). Only relevant if sample_file is specified and is an existing file. In that case, setting append_samples=TRUE will append the samples to the existing file rather than overwriting the contents of the file.

refresh (integer) can be used to control how often the progress of the sampling is reported (i.e. show the progress every refresh iterations). By default, refresh = max(10, iter). The progress indicator is turned off if refresh <= 0.

Deprecated: enable_random_init (logical) being TRUE enables specifying initial values randomly when the initial values are not fully specified from the user.

save_warmup (logical) indicates whether to save draws during the warmup phase and defaults to TRUE. Some memory related problems can be avoided by setting it to FALSE, but some diagnostics are more limited if the warmup draws are not stored.

boost_lib The path for an alternative version of the Boost C++ to use instead of the one in the BH package.
eigen_lib The path for an alternative version of the Eigen C++ library to the one in RcppEigen.

Details

The stan function does all of the work of fitting a Stan model and returning the results as an instance of stanfit. The steps are roughly as follows:

1. Translate the Stan model to C++ code. (stanc)
2. Compile the C++ code into a binary shared object, which is loaded into the current R session (an object of S4 class stanmodel is created). (stan_model)
3. Draw samples and wrap them in an object of S4 class stanfit. (sampling)
The returned object can be used with methods such as `print`, `summary`, and `plot` to inspect and retrieve the results of the fitted model.

`stan` can also be used to sample again from a fitted model under different settings (e.g., different `iter`, `data`, etc.) by using the `fit` argument to specify an existing `stanfit` object. In this case, the compiled C++ code for the model is reused.

**Value**

An object of S4 class `stanfit`. However, if `cores > 1` and there is an error for any of the chains, then the error(s) are printed. If all chains have errors and an error occurs before or during sampling, the returned object does not contain samples. But the compiled binary object for the model is still included, so we can reuse the returned object for another sampling.

**Passing data to Stan**

The data passed to `stan` are preprocessed before being passed to Stan. If data is not a character vector, the data block of the Stan program is parsed and R objects of the same name are searched starting from the calling environment. Then, if data is list-like but not a `data.frame` the elements of data take precedence. This behavior is similar to how a `formula` is evaluated by the `lm` function when data is supplied. In general, each R object being passed to Stan should be either a numeric vector (including the special case of a `scalar`) or a numeric array (matrix). The first exception is that an element can be a logical vector: `TRUE`’s are converted to 1 and `FALSE`’s to 0. An element can also be a data frame or a specially structured list (see details below), both of which will be converted into arrays in the preprocessing. Using a specially structured list is not encouraged though it might be convenient sometimes; and when in doubt, just use arrays.

This preprocessing for each element mainly includes the following:

1. Change the data of type from `double` to `integer` if no accuracy is lost. The main reason is that by default, R uses `double` as data type such as in a `<- 3`. But Stan will not read data of type `int` from `real` and it reads data from `int` if the data type is declared as `real`.
2. Check if there is `NA` in the data. Unlike BUGS, Stan does not allow missing data. Any `NA` values in supplied data will cause the function to stop and report an error.
3. Check data types. Stan allows only numeric data, that is, doubles, integers, and arrays of these. Data of other types (for example, characters and factors) are not passed to Stan.
4. Check whether there are objects in the data list with duplicated names. Duplicated names, if found, will cause the function to stop and report an error.
5. Check whether the names of objects in the data list are legal Stan names. If illegal names are found, it will stop and report an error. See (Cmd)Stan’s manual for the rules of variable names.
6. When an element is of type `data.frame`, it will be converted to `matrix` by function `data.matrix`.
7. When an element is of type `list`, it is supposed to make it easier to pass data for those declared in Stan code such as "vector[J] y1[I]" and "matrix[J,K] y2[I]". Using the latter as an example, we can use a list for `y2` if the list has "I" elements, each of which is an array (matrix) of dimension "J*K". However, it is not possible to pass a list for data declared such as "vector[K] y3[I,J]"; the only way for it is to use an array with dimension "I*J*K". In addition, technically a `data.frame` in R is also a list, but it should not be used for the purpose here since a `data.frame` will be converted to a matrix as described above.
Stan treats a vector of length 1 in R as a scalar. So technically if, for example, "real y[1];" is defined in the data block, an array such as "y = array(1.0, dim = 1)" in R should be used. This is also the case for specifying initial values since the same underlying approach for reading data from R in Stan is used, in which vector of length 1 is treated as a scalar.

In general, the higher the optimization level is set, the faster the generated binary code for the model runs, which can be set in a Makevars file. However, the binary code generated for the model runs fast by using a higher optimization level at the cost of longer times to compile the C++ code.

References


See Also

- The package vignettes for an example of fitting a model and accessing the contents of stanfit objects (http://mc-stan.org/rstan/articles/).
- *stanc* for translating model code in Stan modeling language to C++, *sampling* for sampling, and *stanfit* for the fitted results.
- *as.array.stanfit* and *extract* for extracting samples from stanfit objects.

Examples

```r
## Not run:
#### example 1
library(rstan)
scode <- "
parameters {
  real y[2];
}
model {
  y[1] ~ normal(0, 1);
  y[2] ~ double_exponential(0, 2);
}
" 
fit1 <- stan(model_code = scode, iter = 10, verbose = FALSE)
print(fit1)
fit2 <- stan(fit = fit1, iter = 10000, verbose = FALSE)

## using as.array on the stanfit object to get samples
a2 <- as.array(fit2)

## extract samples as a list of arrays
e2 <- extract(fit2, permuted = FALSE)

#### example 2
#### the result of this package is included in the package
excode <- '
transformed data {
  real y[20];
  y[1] = 0.5796; y[2] = 0.2276; y[3] = -0.2959;
  y[10] = 0.3746; y[11] = 0.4773; y[12] = 0.1803;
  y[16] = 0.9459; y[17] = -0.382; y[18] = 0.7619;
  y[19] = 0.1006; y[20] = -1.7461;
}
parameters {
  real mu;
  real<lower=0, upper=10> sigma;
  vector[2] z[3];
  real<lower=0> alpha;
}
model {
  y ~ normal(mu, sigma);
  for (i in 1:3)
    z[i] ~ normal(0, 1);
  alpha ~ exponential(2);
}

exfit <- stan(model_code = excode, save_dso = FALSE, iter = 500)
print(exfit)
plot(exfit)

## End(Not run)
## Not run:
## examples of specify argument 'init' for function stan

## define a function to generate initial values that can
## be fed to function stan's argument 'init'
# function form 1 without arguments
initf1 <- function() {
  list(mu = 1, sigma = 4, z = array(rnorm(6), dim = c(3,2)), alpha = 1)
}
# function form 2 with an argument named 'chain_id'
initf2 <- function(chain_id = 1) {
  # cat("chain_id =", chain_id, "\n")
  list(mu = 1, sigma = 4, z = array(rnorm(6), dim = c(3,2)), alpha = chain_id)
}

# generate a list of lists to specify initial values
n_chains <- 4
init_ll <- lapply(1:n_chains, function(id) initf2(chain_id = id))

exfit0 <- stan(model_code = excode, init = initf1)
stan(fit = exfit0, init = initf2)
stan(fit = exfit0, init = init_ll, chains = n_chains)

## End(Not run)
Translate Stan model specification to C++ code

**Description**

Translate a model specification in Stan code to C++ code, which can then be compiled and loaded for sampling.

**Usage**

```r
stanc(file, model_code = '', model_name = "anon_model", verbose = FALSE,
    obfuscate_model_name = TRUE, allow_undefined = FALSE,
    isystem = c(if (!missing(file)) dirname(file), getwd()))
stanc_builder(file, isystem = c(dirname(file), getwd()),
    verbose = FALSE, obfuscate_model_name = FALSE,
    allow_undefined = FALSE)
```

**Arguments**

- **file**: A character string or a connection that R supports specifying the Stan model specification in Stan's modeling language.
- **model_code**: Either a character string containing a Stan model specification or the name of a character string object in the workspace. This parameter is used only if parameter `file` is not specified, so it defaults to the empty string.
- **model_name**: A character string naming the model. The default is "anon_model". However, the model name will be derived from `file` or `model_code` (if `model_code` is the name of a character string object) if `model_name` is not specified.
- **verbose**: Logical, defaulting to FALSE. If TRUE more intermediate information is printed during the translation procedure.
- **obfuscate_model_name**: Logical, defaulting to TRUE, indicating whether to use a randomly-generated character string for the name of the C++ class. This prevents name clashes when compiling multiple models in the same R session.
- **isystem**: A character vector naming a path to look for file paths in `file` that are to be included within the Stan program named by `file`. See the Details section below.
- **allow_undefined**: A logical scalar defaulting to FALSE indicating whether to allow Stan functions to be declared but not defined in `file` or `model_code`. If TRUE, then it is the caller's responsibility to provide a function definition in another header file or linked shared object.

**Details**

The `stanc_builder` function supports the standard C++ convention of specifying something like `#include "my_includes.txt"` on an entire line within the file named by the `file` argument.
In other words, `stanc_builder` would look for "my_includes.txt" in (or under) the directories
named by the `isystem` argument and — if found — insert its contents verbatim at that position
before calling `stanc` on the resulting `model_code`. This mechanism reduces the need to copy
common chunks of code across Stan programs. It is possible to include such files recursively.

Note that line numbers referred to in parser warnings or errors refer to the postprocessed Stan
program rather than file. In the case of a parser error, the postprocessed Stan program will be
printed after the error message. Line numbers referred to in messages while Stan is executing also
refer to the postprocessed Stan program which can be obtained by calling `get_stancode`.

Value

A list with named entries:

1. `model_name` Character string for the model name.
2. `model_code` Character string for the model’s Stan specification.
3. `cppcode` Character string for the model’s C++ code.
4. `status` Logical indicating success/failure (always `TRUE`) of translating the Stan code.

Note

Unlike R, in which variable identifiers may contain dots (e.g. a\_1), Stan prohibits dots from occurring
in variable identifiers. Furthermore, C++ reserved words and Stan reserved words may not be
used for variable names; see the Stan User’s Guide for a complete list.

References


See Also

`stan_model` and `stan`.

Examples

```r
stanmodelcode <- "
data {
  int<lower=0> N;
  real y[N];
}

parameters {
  real mu;
}

model {
  mu ~ normal(0, 1);
  y ~ normal(mu, 1);
}"
```
Description

The components (slots) of a stanfit object and the various available methods are described below. When methods have their own more detailed documentation pages links are provided.

Objects from the Class

An object of class stanfit contains the output derived from fitting a Stan model as returned by the top-level function stan or the lower-level methods sampling and vb (which are defined on class stanmodel). Many methods (e.g., print, plot, summary) are provided for summarizing results and various access methods also allow the underlying data (e.g., simulations, diagnostics) contained in the object to be retrieved.

Slots

model_name: The model name as a string.

model_pars: A character vector of names of parameters (including transformed parameters and derived quantities).

par_dims: A named list giving the dimensions for all parameters. The dimension for a scalar parameter is given as numeric(0).

mode: An integer indicating the mode of the fitted model. 0 indicates sampling mode, 1 indicates test gradient mode (no sampling is done), and 2 indicates error mode (an error occurred before sampling). Most methods for stanfit objects are useful only if mode=0.

sim: A list containing simulation results including the posterior draws as well as various pieces of metadata used by many of the methods for stanfit objects.

inits: The initial values (either user-specified or generated randomly) for all chains. This is a list with one component per chain. Each component is a named list containing the initial values for each parameter for the corresponding chain.

stan_args: A list with one component per chain containing the arguments used for sampling (e.g. iter, seed, etc.).

stanmodel: The instance of S4 class stanmodel.

date: A string containing the date and time the object was created.

.MISC: Miscellaneous helper information used for the fitted model. This is an object of type environment. Users rarely (if ever) need to access the contents of .MISC.
Methods

Printing, plotting, and summarizing:

show Print the default summary for the model.
print Print a customizable summary for the model. See print.stanfit.
plot Create various plots summarizing the fitted model. See plot.stanfit-method.
summary Summarize the distributions of estimated parameters and derived quantities using the posterior draws. See summary.stanfit-method.
get_posterior_mean Get the posterior mean for parameters of interest (using pars to specify a subset of parameters). Returned is a matrix with one column per chain and an additional column for all chains combined.

Extracting posterior draws:
extract Extract the draws for all chains for all (or specified) parameters. See extract.
as.array, as.matrix, as.data.frame Coerce the draws (without warmup) to an array, matrix or data frame. See as.array.stanfit.
As.mcmc.list Convert a stanfit object to an mcmc.list as in package coda. See As.mcmc.list.
get_logposterior Get the log-posterior at each iteration. Each element of the returned list is the vector of log-posterior values (up to an additive constant, i.e. up to a multiplicative constant on the linear scale) for a single chain. The optional argument inc_warmup (defaulting to TRUE) indicates whether to include the warmup period.

Diagnostics, log probability, and gradients:
get_sampler_params Obtain the parameters used for the sampler such as stepsize and treedepth. The results are returned as a list with one component (an array) per chain. The array has number of columns corresponding to the number of parameters used in the sampler and its column names provide the parameter names. Optional argument inc_warmup (defaulting to TRUE) indicates whether to include the warmup period.
get_adaptation_info Obtain the adaptation information for the sampler if NUTS was used. The results are returned as a list, each element of which is a character string with the info for a single chain.

log_prob Compute the log probability density (lp__) for a set of parameter values (on the unconstrained space) up to an additive constant. The unconstrained parameters are specified using a numeric vector. The number of parameters on the unconstrained space can be obtained using method get_num_upars. A numeric value is returned. See also the documentation in log_prob.

grad_log_prob Compute the gradient of log probability density function for a set of parameter values (on the unconstrained space) up to an additive constant. The unconstrained parameters are specified using a numeric vector with the length being the number of unconstrained parameters. A numeric vector is returned with the length of the number of unconstrained parameters and an attribute named log_prob being the lp__. See also the documentation in grad_log_prob.

get_num_upars Get the number of unconstrained parameters of the model. The number of parameters for a model is not necessarily equal to this number of unconstrained parameters. For example, when a parameter is specified as a simplex of length K, the number of unconstrained parameters is K-1.
unconstrain_pars  Transform the parameters to unconstrained space. The input is a named list as for specifying initial values for each parameter. A numeric vector is returned. See also the documentation in unconstrain_pars.

constrain_pars  Get the parameter values from their unconstrained space. The input is a numeric vector. A list is returned. This function is contrary to unconstrain_pars. See also the documentation in constrain_pars.

**Metadata and miscellaneous:**

get_stancode  Get the Stan code for the fitted model as a string. The result can be printed in a readable format using cat.

get_stanmodel  Get the object of S4 class stanmodel of the fitted model.

get_elapsed_time  Get the warmup time and sample time in seconds. A matrix of two columns is returned with each row containing the warmup and sample times for one chain.

get_inits, iter = NULL  Get the initial values for parameters used in sampling all chains. The returned object is a list with the same structure as the inits slot described above. If object@mode=2 (error mode) an empty list is returned. If iter is not NULL, then the draw from that iteration is returned for each chain rather than the initial state.

get_cppmode  Get the optimization mode used for compilation. The returned string is one of "fast", "presentation2", "presentation1", and "debug".

get_seed  Get the (P)RNG seed used. When the fitted object is empty (mode=2), NULL might be returned. In the case that the seeds for all chains are different, use get_seeds.

get_seeds  Get the seeds used for all chains. When the fitted object is empty (mode=2), NULL might be returned.

**References**


**See Also**

stan and stanmodel

**Examples**

```r
## Not run:
showClass("stanfit")
ecode <- '
parameters {
  real<lower=0> y[2];
}
model {
  y ~ exponential(1);
}
',
fit <- stan(model_code = ecode, iter = 10, chains = 1)
fit2 <- stan(fit = fit)
print(fit2)
```
```r
plot(fit2)
traceplot(fit2)
ainfo <- get_adaptation_info(fit2)
cat(ainfo[[1]])
seed <- get_seed(fit2)
sp <- get_sampler_params(fit2)
spR <- get_sampler_params(fit2, inc_warmup = FALSE)
head(sp[[1]])

lp <- log_prob(fit, c(1, 2))
grad <- grad_log_prob(fit, c(1, 2))
lpR <- attr(grad, "log_prob") # should be the same as "lp"

# get the number of parameters on the unconstrained space
n <- get_num_upars(fit)

# parameters on the positive real line (constrained space)
y1 <- list(y = rep(1, 2))

uy <- unconstrain_pars(fit, y1)
## uy should be c(0, 0) since here the log transformation is used
y1star <- constrain_pars(fit, uy)

print(y1)
print(y1star) # y1star should equal to y1

## End(Not run)

# Create a stanfit object from reading CSV files of samples (saved in rstan package) generated by function stan for demonstration purpose from model as follows.

```
```stan
transformed data {
    real y[20];
    y[1] <- 0.5796; y[2] <- 0.2276; y[3] <- -0.2959;
    y[13] <- 0.5215; y[14] <- -1.6044; y[15] <- -0.6783;
    y[16] <- 0.9459; y[17] <- -0.382; y[18] <- 0.7619;
    y[19] <- 0.1006; y[20] <- -1.7461;
}
parameters {
    real mu;
    real<lower=0, upper=10> sigma;
    vector[2] z[3];
    real<lower=0> alpha;
}
model {
    y ~ normal(mu, sigma);
    for (i in 1:3)
        z[i] ~ normal(0, 1);
    alpha ~ exponential(2);
```
stanmodel-class

Class representing model compiled from C++

Description

A stanmodel object represents the model compiled from C++ code. The sampling method defined in this class may be used to draw samples from the model and optimizing method is for obtaining a point estimate by maximizing the log-posterior.

Objects from the Class

Instances of stanmodel are usually created by calling function stan_model or function stan.

Slots

- `model_name`: The model name, an object of type character.
- `model_code`: The Stan model specification, an object of type character.
- `model_cpp`: Object of type list that includes the C++ code for the model.
- `mk_cppmodule`: A function to return a RCpp module. This function will be called in function `sampling` and `optimizing` with one argument (the instance of stanmodel itself).
- `dso`: Object of S4 class cxxdso. The container for the dynamic shared objects compiled from the C++ code of the model, returned from function `cxxfunction` in package inline.
Methods

show signature(object = "stanmodel"): print the Stan model specification.

vb signature(object = "stanmodel"): use the variational Bayes algorithms.

sampling signature(object = "stanmodel"): draw samples for the model (see sampling).

optimizing signature(object = "stanmodel"): obtain a point estimate by maximizing the posterior (see optimizing).

get_cppcode signature(object = "stanmodel"): returns the C++ code for the model as a character string. This is part of the C++ code that is compiled to the dynamic shared object for the model.

get_stancode signature(object = "stanmodel"): returns the Stan code for the model as a character string.

get_cxxflags signature(object = "stanmodel"): return the cxxflags used for compiling the model. The returned string is like CXXFLAGS = -O3.

Note

Objects of class stanmodel can be saved for use across R sessions only if save_dso = TRUE is set during calling functions that create stanmodel objects (e.g., stan and stan_model).

Even if save_dso = TRUE, the model cannot be loaded on a platform (operating system, 32 bits or 64 bits, etc.) that differs from the one on which it was compiled.

See Also

stan_model, stanc, sampling, optimizing, vb

Examples

showClass("stanmodel")

Description

Stan includes a variety of examples and most of the BUGS example models that are translated into Stan modeling language. One example is chosen from a list created from matching user input and gets fitted in the demonstration.

Usage

stan_demo(model = character(0),
          method = c("sampling", "optimizing", "meanfield", "fullrank"), ...)

Demonstrate examples included in Stan
Arguments

model
A character string for model name to specify which model will be used for demonstration. The default is an empty string, which prompts the user to select one the available models. If \texttt{model = 0}, a character vector of all models is returned without any user intervention. If \texttt{model = i} where \texttt{i > 0}, then the ith available model is chosen without user intervention, which is useful for testing.

method
Whether to call \texttt{sampling} (the default), \texttt{optimizing}, or one of the variants of \texttt{vb} for the demonstration

...Further arguments passed to method.

Value

An S4 object of \texttt{stanfit}, unless \texttt{model = 0}, in which case a character vector of paths to available models is returned.

References


See Also

\texttt{sampling,optimizing}

Examples

## Not run:
    dogsfit <- \texttt{stan_demo("dogs")} # run the dogs model
    fit1 <- \texttt{stan_demo(1)} # run model_names[1]

## End(Not run)

---

\textit{stan\_model} \hspace{1cm} \textit{Construct a Stan model}

Description

Construct an instance of S4 class \texttt{stanmodel} from a model specified in Stan’s modeling language. A \texttt{stanmodel} object can then be used to draw samples from the model. The Stan program (the model expressed in the Stan modeling language) is first translated to C++ code and then the C++ code for the model plus other auxiliary code is compiled into a dynamic shared object (DSO) and then loaded. The loaded DSO for the model can be executed to draw samples, allowing inference to be performed for the model and data.
stan_model

Usage

stan_model(file, model_name = "anon_model",
model_code = "", stanc_ret = NULL,
boost_lib = NULL, eigen_lib = NULL,
save_dso = TRUE, verbose = FALSE,
auto_write = rstan_options("auto_write"),
obfuscate_model_name = TRUE,
allow_undefined = FALSE, includes = NULL,
isystem = c(if (!missing(file)) dirname(file), getwd())))

Arguments

file A character string or a connection that R supports specifying the Stan model specification in Stan’s modeling language.
model_name A character string naming the model; defaults to “anon_model”. However, the model name will be derived from file or model_code (if model_code is the name of a character string object) if model_name is not specified.
model_code Either a character string containing the model specification or the name of a character string object in the workspace. This is an alternative to specifying the model via the file or stanc_ret arguments.
stanc_ret A named list returned from a previous call to the stanc function. The list can be used to specify the model instead of using the file or model_code arguments.
boost_lib The path to a version of the Boost C++ library to use instead of the one in the BH package.
eigen_lib The path to a version of the Eigen C++ library to use instead of the one in the RcppEigen package.
save_dso Logical, defaulting to TRUE, indicating whether the dynamic shared object (DSO) compiled from the C++ code for the model will be saved or not. If TRUE, we can draw samples from the same model in another R session using the saved DSO (i.e., without compiling the C++ code again).
verbose Logical, defaulting to FALSE, indicating whether to report additional intermediate output to the console, which might be helpful for debugging.
auto_write Logical, defaulting to the value of rstan_options("auto_write"), indicating whether to write the object to the hard disk using saveRDS. Although this argument is FALSE by default, we recommend calling rstan_options("auto_write" = TRUE) in order to avoid unnecessary recompilations. If file is supplied and its dirname is writable, then the object will be written to that same directory, substituting a .rds extension for the .stan extension. Otherwise, the object will be written to the tempdir.
obfuscate_model_name A logical scalar that is TRUE by default and passed to stanc.
allow_undefined A logical scalar that is FALSE by default and passed to stanc.
includes If not NULL (the default), then a character vector of length one (possibly containing one or more "\n") of the form '#include "/full/path/to/my_header.hpp"',
which will be inserted into the C++ code in the model’s namespace and can be used to provide definitions of functions that are declared but not defined in file or model_code when allow_undefined = TRUE

isystem A character vector naming a path to look for file paths in file that are to be included within the Stan program named by file. See the Details section below.

Details

If a previously compiled stanmodel exists on the hard drive, its validity is checked and then returned without recompiling. The most common form of invalidity seems to be Stan code that ends with a } rather than a blank line, which causes the hash checker to think that the current model is different than the one saved on the hard drive. To avoid reading previously compiled stanmodels from the hard drive, supply the stanc_ret argument rather than the file or model_code arguments.

There are three ways to specify the model’s code for stan_model:

1. parameter model_code: a character string containing the Stan model specification,
2. parameter file: a file name (or a connection) from which to read the Stan model specification, or
3. parameter stanc_ret: a list returned by stanc to be reused.

Value

An instance of S4 class stanmodel that can be passed to the sampling, optimizing, and vb functions.

References


See Also

stanmodel for details on the class.

sampling, optimizing, and vb, which take a stanmodel object as input, for estimating the model parameters.

More details on Stan, including the full user’s guide and reference manual, can be found at http://mc-stan.org/.

Examples

```r
## Not run:
stancode <- 'data {real y_mean;} parameters {real y;} model {y ~ normal(y_mean,1);}'
mod <- stan_model(model_code = stancode, verbose = TRUE)
fit <- sampling(mod, data = list(y_mean = 0))
fit2 <- sampling(mod, data = list(y_mean = 5))

## End(Not run)
```
**Description**

This function takes a vector of names of \( \mathcal{R} \) objects and outputs text representations of the objects to a file or connection. The file created by `stan_rdump` is typically used as data input of the Stan package ([http://mc-stan.org/](http://mc-stan.org/)) or `sourced` into another \( \mathcal{R} \) session. The usage of this function is very similar to `dump` in \( \mathcal{R} \).

**Usage**

```r
stan_rdump(list, file = "", append = FALSE,
    envir = parent.frame(),
    width = options("width")$width,
    quiet = FALSE)
```

**Arguments**

- `list`  
  A vector of character string: the names of one or more \( \mathcal{R} \) objects to be dumped. See the note below.

- `file`  
  Either a character string naming a file or a `connection`. "" indicates output to the console.

- `append`  
  Logical: if TRUE and file is a character string, output will be appended to file; otherwise, it will overwrite the contents of file.

- `envir`  
  The environment to search for objects.

- `width`  
  The width for maximum characters on a line. The output is broken into lines with width.

- `quiet`  
  Whether to suppress warning messages that would appear when a variable is not found or not supported for dumping (not being numeric or it would not be converted to numeric) or a variable name is not allowed in Stan.

**Value**

An invisible character vector containing the names of the objects that were dumped.

**Note**

`stan_rdump` only dumps numeric data, which first can be a scalar, vector, matrix, or (multidimensional) array. Additional types supported are logical (TRUE and FALSE), factor, data.frame and a specially structured list.

The conversion for logical variables is to map TRUE to 1 and FALSE to 0. For factor variable, function `as.integer` is used to do the conversion (If we want to transform a factor f to approximately its original numeric values, see the help of function factor and do the transformation before calling `stan_rdump`). In the case of data.frame, function `data.matrix` is applied to the data frame before
dumping. See the notes in stan for the specially structured list, which will be converted to array before dumping.

stan_rdump will check whether the names of objects are legal variable names in Stan. If an illegal name is found, data will be dumped with a warning. However, passing the name checking does not necessarily mean that the name is legal. More details regarding rules of variable names in Stan can be found in Stan’s manual.

If objects with specified names are not found, a warning will be issued.

References


See Also
dump

Examples

# set variables in global environment
a <- 17.5
b <- c(1,2,3)
# write variables a and b to file ab.data.R in temporary directory
stan_rdump(c('a','b'), file.path(tempdir(), "ab.data.R"))
unlink(file.path(tempdir(), "ab.data.R"))

x <- 1; y <- 1:10; z <- array(1:10, dim = c(2,5))
stan_rdump(ls(pattern = "^[xyz]"), file.path(tempdir(), "xyz.Rdump"))
cat(paste(readLines(file.path(tempdir(), "xyz.Rdump")), collapse = '\n'), '\n')
unlink(file.path(tempdir(), "xyz.Rdump"))

---

**stan_version**

*Obtain the version of Stan*

**Description**

The stan version is in form of major.minor.patch; the first version is 1.0.0, indicating major version 1, minor version 0 and patch level 0. Functionality only changes with minor versions and backward compatibility will only be affected by major versions.

**Usage**

stan_version()

**Value**

A character string giving the version of Stan used in this version of RStan.
References


See Also

stan and stan_model

Examples

stan_version()

summary-methods  Summary method for stanfit objects

Description

Summarize the distributions of estimated parameters and derived quantities using the posterior draws.

Usage

## S4 method for signature 'stanfit'
summary(object, pars, probs = c(0.025, 0.25, 0.50, 0.75, 0.975),
use_cache = TRUE, ...)

Arguments

object  An instance of class stanfit.
pars  A character vector of parameter names. Defaults to all parameters as well as the log-posterior (lp__).
probs  A numeric vector of quantiles of interest. The default is c(0.025, 0.25, 0.5, 0.75, 0.975).
use_cache  Logical, defaulting to TRUE. When use_cache=TRUE the summary quantities for all parameters are computed and cached for future use. Setting use_cache=FALSE can be used to avoid performing the summary computations for all parameters if pars is given as some specific parameters.
...  Currently unused.

Value

The summary method returns a named list with elements summary and c_summary, which contain summaries for for all chains merged and individual chains, respectively. Included in the summaries are quantiles, means, standard deviations (sd), effective sample sizes (n_eff), and split Rhat (the potential scale reduction derived from all chains after splitting each chain in half and treating the halves as chains). For the summary of all chains merged, Monte Carlo standard errors (se_mean) are also reported.
See Also

- `monitor`, which computes similar summaries but accepts an array of MCMC draws as its input rather than a `stanfit` object.
- The RStan vignettes for more example usage.

Examples

```r
## Not run:
ecode <- 'parameters {
   real<lower=0> y[2];
}
model {
   y ~ exponential(1);
}
fit <- stan(model_code = ecode)
s <- summary(fit, probs = c(0.1, 0.9))
s$s$summary # all chains merged
s$c_summary # individual chains

## End(Not run)
```

---

**traceplot**

*Markov chain traceplots*

**Description**

Draw the traceplot corresponding to one or more Markov chains, providing a visual way to inspect sampling behavior and assess mixing across chains and convergence.

**Usage**

```r
## S4 method for signature 'stanfit'
traceplot(object, pars, include = TRUE, unconstrain = FALSE,
          inc_warmup = FALSE, window = NULL, nrow = NULL, ncol = NULL, ...)
```

**Arguments**

- `object` An instance of class `stanfit`.
- `pars` A character vector of parameter names. Defaults to all parameters or the first 10 parameters (if there are more than 10).
- `include` Should the parameters given by the `pars` argument be included (the default) or excluded from the plot? Only relevant if `pars` is not missing.
- `inc_warmup` TRUE or FALSE, indicating whether the warmup sample are included in the trace plot; defaults to FALSE.
traceplot

window
A vector of length 2. Iterations between window[1] and window[2] will be shown in the plot. The default is to show all iterations if inc_warmup is TRUE and all iterations from the sampling period only if inc_warmup is FALSE. If inc_warmup is FALSE the iterations specified in window should not include iterations from the warmup period.

unconstrain
Should parameters be plotted on the unconstrained space? Defaults to FALSE.

nrow,ncol
Passed to facet_wrap.

... Optional arguments to pass to geom_path (e.g. size, linetype, alpha, etc.).

Value
A ggplot object that can be further customized using the ggplot2 package.

Methods

traceplot signature(object = "stanfit") Plot the sampling paths for all chains.

See Also

List of RStan plotting functions, Plot options

Examples

```r
# Not run:
# Create a stanfit object from reading CSV files of samples (saved in rstan
# package) generated by function stan for demonstration purpose from model as follows.
#
# excode <- 'transformed data {
#    real y[20];
#    y[1] <- 0.5796; y[2] <- 0.2276; y[3] <- -0.2959;
#    y[13] <- 0.5215; y[14] <- -1.6044; y[15] <- -0.6703;
#    y[16] <- 0.9459; y[17] <- 0.3802; y[18] <- 0.7619;
#    y[19] <- 0.1006; y[20] <- -1.7461;
#}
parameters {
    real mu;
    real<lower=0, upper=10> sigma;
    vector[2] z[3];
    real<lower=0> alpha;
}
model {
    y ~ normal(mu, sigma);
    for (i in 1:3)
        z[i] ~ normal(0, 1);
    alpha ~ exponential(2);
}
```
# exfit <- stan(model_code = excode, save_dso = FALSE, iter = 200,  
#     sample_file = "rstan_doc_ex.csv")  
# exfit <- read_stan_csv(dir(system.file('misc', package = 'rstan'),  
#     pattern='rstan_doc_ex_[[:digit:]].csv',  
#     full.names = TRUE))

print(exfit)  
traceplot(exfit)  
traceplot(exfit, size = 0.25)  
traceplot(exfit, pars = "sigma", inc_warmup = TRUE)  

trace <- traceplot(exfit, pars = c("z[1,1]", "z[3,1]"))  
trace + scale_color_discrete() + theme(legend.position = "top")  

## End(Not run)

---

**Run Stan’s variational algorithm for approximate posterior sampling**

**Description**

Approximately draw from a posterior distribution using variational inference.

This is still considered an experimental feature. We recommend calling `stan` or `sampling` for final inferences and only using `vb` to get a rough idea of the parameter distributions.

**Usage**

```r
## S4 method for signature 'stanmodel'
vb(object, data = list(), pars = NA, include = TRUE,  
    seed = sample.int(.Machine$integer.max, 1),  
    init = 'random', check_data = TRUE,  
    sample_file = tempfile(fileext = '.csv'),  
    algorithm = c("meanfield", "fullrank"),  
    importance_resampling = FALSE, keep_every = 1,  
    ...)```

**Arguments**

- `object` An object of class `stanmodel`.
- `data` A named list or environment providing the data for the model or a character vector for all the names of objects used as data. See the **Passing data to Stan** section in `stan`.
- `pars` If not NA, then a character vector naming parameters, which are included in the output if `include = TRUE` and excluded if `include = FALSE`. By default, all parameters are included.
include Logical scalar defaulting to TRUE indicating whether to include or exclude the parameters given by the pars argument. If FALSE, only entire multidimensional parameters can be excluded, rather than particular elements of them.

seed The seed for random number generation. The default is generated from 1 to the maximum integer supported by R on the machine. Even if multiple chains are used, only one seed is needed, with other chains having seeds derived from that of the first chain to avoid dependent samples. When a seed is specified by a number, as.integer will be applied to it. If as.integer produces NA, the seed is generated randomly. The seed can also be specified as a character string of digits, such as "12345", which is converted to integer.

init Initial values specification. See the detailed documentation for the init argument in stan.

check_data Logical, defaulting to TRUE. If TRUE the data will be preprocessed; otherwise not. See the Passing data to Stan section in stan.

sample_file A character string of file name for specifying where to write samples for all parameters and other saved quantities. This defaults to a temporary file.

algorithm Either "meanfield" (the default) or "fullrank", indicating which variational inference algorithm is used. The "meanfield" option uses a fully factorized Gaussian for the approximation whereas the fullrank option uses a Gaussian with a full-rank covariance matrix for the approximation. Details and additional references are available in the Stan manual.

importance_resampling Logical scalar (defaulting to FALSE) indicating whether to do importance resampling to adjust the draws at the optimum to be more like draws from the posterior distribution

keep_every Integer scalar (defaulting to 1) indicating the interval by which to thin the draws when importance_resampling = TRUE

... Other optional parameters:

- iter (positive integer), the maximum number of iterations, defaulting to 10000.
- grad_samples (positive integer), the number of samples for Monte Carlo estimate of gradients, defaulting to 1.
- elbo_samples (positive integer), the number of samples for Monte Carlo estimate of ELBO (objective function), defaulting to 100. (ELBO stands for "the evidence lower bound").
- eta (double), positive stepsize weighting parameter for variational inference but is ignored if adaptation is engaged, which is the case by default.
- adapt_engaged (logical), a flag indicating whether to automatically adapt the stepsize, defaulting to TRUE.
- tol_rel_obj (positive double), the convergence tolerance on the relative norm of the objective, defaulting to 0.01.
- eval_elbo (positive integer), evaluate ELBO every Nth iteration, defaulting to 100.
- output_samples (positive integer), number of posterior samples to draw and save, defaults to 1000.
• adapt_iter (positive integer), the maximum number of iterations to adapt the stepsize, defaulting to 50. Ignored if adapt_engaged = FALSE.

Refer to the manuals for both CmdStan and Stan for more details.

Value

An object of stanfit-class.

Methods

vb signature(object = "stanmodel")

Call Stan’s variational Bayes methods for the model defined by S4 class stanmodel given the data, initial values, etc.

References


See Also

stanmodel

The manuals of CmdStan and Stan.

Examples

```r
## Not run:
m <- stan_model(model_code = 'parameters (real y;) model (y ~ normal(0,1);)')
f <- vb(m)

## End(Not run)
```
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